

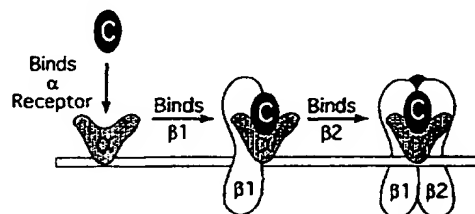


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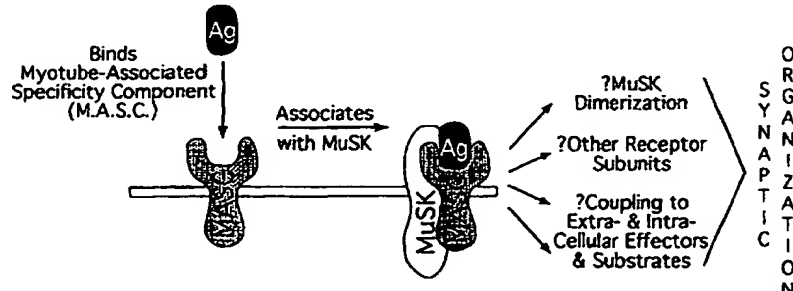
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<p>(21) International Application Number: PCT/US96/20696</p> <p>(22) International Filing Date: 13 December 1996 (13.12.96)</p> <p>(30) Priority Data: 60/008,657 15 December 1995 (15.12.95) US 08/644,271 10 May 1996 (10.05.96) US</p> <p>(71) Applicant (for all designated States except US): REGENERON PHARMACEUTICALS, INC. [US/US]; 777 Old Saw Mill River Road, Tarrytown, NY 10591-6707 (US).</p> <p>(72) Inventors; and</p> <p>(75) Inventors/Applicants (for US only): VALENZUELA, David, M. [CL/US]; 216 Grange Street, Franklin Square, NY 11010 (US). GLASS, David, J. [US/US]; 341 Furnace Dock Road, Cortlandt Manor, NY 10566 (US). BOWEN, David, C. [US/US]; 679 Warburton Avenue, Yonkers, NY 10701 (US). YANCOPOULOS, George, D. [US/US]; 1519 Baptist Church Road, Yorktown Heights, NY 10598 (US).</p> <p>(74) Agents: COBERT, Robert, J.; Regeneron Pharmaceuticals, Inc., 777 Old Saw Mill River Road, Tarrytown, NY 10591 (US) et al.</p>		<p>(81) Designated States: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, US, UZ, VN, ARIPO patent (KE, LS, MW, SD, SZ, UG), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG).</p> <p>Published Without international search report and to be republished upon receipt of that report.</p>

(54) Title: NOVEL TYROSINE KINASE RECEPTORS AND LIGANDS

A. Formation of CNTF Receptor Complex

B. CNTF Receptor Complex With Soluble β Components

C. Formation of Receptor Complex For Agrin



(57) Abstract

The present invention provides for a gene, designated as *muskl*, that encodes a novel tyrosine kinase receptor expressed in high levels in denervated muscle. The invention also provides for an isolated polypeptide which activates MuSK receptor. The invention further provides for a polypeptide which is functionally equivalent to the MuSK activating polypeptide. The invention also provides assay systems that may be used to detect and/or measure ligands that bind the *muskl* gene product. The present invention also provides for diagnostic and therapeutic methods based on molecules that activate MuSK.

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NOVEL TYROSINE KINASE RECEPTORS AND LIGANDS

This application claims priority of United States Application Serial No. 08/644,271 filed May 10, 1996 and of United States Provisional Application No. 60/008,657 filed December 15, 1995, each of which is incorporated by reference herein.

Throughout this application various publications are referenced. The disclosures of these publications in their entireties are hereby incorporated by reference into this application.

INTRODUCTION

The present invention provides for a novel receptor molecule, a novel molecule capable of activating the receptor, and methods of making and use thereof.

BACKGROUND OF THE INVENTION

The ability of polypeptide ligands to bind cells and thereby elicit a phenotypic response such as cell growth, survival or differentiation in such cells is often mediated through receptor tyrosine kinases. The extracellular portion of each receptor tyrosine kinase (RTK) is generally the most distinctive portion of the molecule, as it provides the protein with its ligand-recognizing characteristic. Binding of a ligand to the extracellular domain results in signal transduction via an intracellular tyrosine kinase catalytic domain which transmits a biological signal to intracellular target proteins. The particular array of sequence motifs of this cytoplasmic, catalytic domain determines its access to potential kinase substrates (Mohammadi, et al., 1990, Mol. Cell. Biol., 11: 5068-5078; Fantl, et al., 1992, Cell, 69:413-413).

The tissue distribution of a particular tyrosine kinase receptor within higher organisms provides relevant data as to the biological function of the receptor. For example, the localization of a Trk family receptor, TrkB, in tissue provided some insight into the potential biological role of this receptor, as well as the ligands that bind this receptor (referred to herein as cognates). Thus, for example, in adult mice, trkB was found to be preferentially expressed in brain tissue, although significant levels of trkB mRNAs were also observed in lung, muscle, and ovaries. Further, trkB transcripts were detected in mid and late gestation embryos. In situ hybridization analysis of 14 and 18 day old mouse embryos indicated that trkB transcripts were localized in the central and peripheral nervous systems, including brain, spinal cord, spinal and cranial ganglia, paravertebral trunk of the sympathetic nervous system and various innervation pathways, suggesting that the trkB gene product may be a receptor involved in neurogenesis and early neural development as well as play a role in the adult nervous system.

The cellular environment in which an RTK is expressed may influence the biological response exhibited upon binding of a ligand to the receptor. Thus, for example, when a neuronal cell expressing a Trk receptor is exposed to a neurotrophin which binds that receptor, neuronal survival and differentiation results. When the same receptor is expressed by a fibroblast, exposure to the neurotrophin results in proliferation of the fibroblast (Glass, et al., 1991, Cell 66:405-413). Thus, it appears that the extracellular domain provides the determining factor as to the ligand specificity, and once signal transduction is initiated the cellular environment will determine the phenotypic outcome of that signal transduction.

A number of RTK families have been identified based on sequence homologies of their intracellular domains. For example, two members of the TIE (tyrosine kinase with immunoglobulin and EGF homology domains) family, known as TIE-1 and TIE-2, have 79% sequence homology in their

intracellular region (Maisonpierre, et al., 1993, Oncogene 8:1631-1637).

Although these receptors share similar motifs in their extracellular domain, only 32% of the sequences are identical.

- 5 A receptor having a kinase domain that is related to the Trk family has been identified in the electric ray Torpedo californica and may play a role in motor neuron induced synapses on muscle fibers. Jennings, et al. Proc. Natl. Acad. Sci. USA 90: 2895-2899 (1993). This kinase was isolated from the electric organ, a tissue which is a specialized form of skeletal muscle. The tyrosine
- 10 kinase domain of this protein is related to the Trk family, while the extracellular domain is somewhat divergent from the Trks. The protein was found to be expressed at high levels in Torpedo skeletal muscle, and at much lower levels in adult Torpedo brain, spinal cord, heart, liver and testis.
- 15 Often such novel RTKs are identified and isolated by searching for additional members of known families of tyrosine kinase receptors using, for example, PCR-based screens involving known regions of homology among Trk family members. (See, for example, Maisonpierre, et al., 1993, Oncogene 8: 1631-1637). Isolation of such so called "orphan" tyrosine kinase receptors, for which no
- 20 ligand is known, and subsequent determination of the tissues in which such receptors are expressed, provides insight into the regulation of the growth, proliferation and regeneration of cells in target tissues. The identification and isolation of novel RTKs may be used as a means of identifying new
- 25 ligands or activating molecules that may then be used to regulate the survival, growth, differentiation and/or regeneration of cells expressing the receptors. Further, because RTKs appear to mediate a number of important functions during development, the identification and isolation of such
- 30 receptors, ligands and activating molecules enhances our understanding of developmental processes and may improve our ability to diagnose or treat abnormal conditions.

For example, the above described methods may be used to study an event that occurs during development of the neuromuscular junction (NMJ) - the localization of acetylcholine receptors at the synapse. It has long been known that important signals are exchanged across the NMJ (Nitkin et al., 1987, J.Cell.Biol. 105: 2471-2478; Hall, Z.W. and Sanes, J.R., 1993, Cell/Neuron (Suppl.) 72/10: 99-121; Bowe, M.A. and Fallon, J.R., 1995, Ann. Rev. Neurosci. 18: 443-462; Sanes, J.R., 1995, Devel. Biol. 6: 163-173; Burden, S.J., et al., 1995, Devel. Biol. 6: 59-65). These signals include the chemical transmitter, acetylcholine, which is released from vesicles in the nerve terminal, recognized by acetylcholine receptors (AChRs) on the muscle, and ultimately results in electrical activation and contraction of the muscle.

Muscle also provides neurotrophic factors that support survival of motor neurons (DeChiara, T. et al., 1995, Cell 83: 313-322), and the nerve may in turn provide myotrophic factors that maintain muscle mass (Helgren, M.E., et al., 1994, Cell 76: 493-504). Reciprocal signaling interactions are also critical both for the formation and maintenance of the neuromuscular junction itself. Such signals regulate recognition of nerve-to-muscle contact, arrest the growth of the incoming nerve ending, and induce formation of a highly specialized nerve terminal marked by a polarized arrangement of synaptic vesicles and active zones. Simultaneously, precisely juxtaposed with respect to the nerve terminal, a complex molecular apparatus forms on the muscle membrane. This specialized postsynaptic structure, termed the motor endplate, comprises a tiny patch on the muscle membrane which is characterized by a dense clustering of particular proteins; some of these may receive nerve-derived signals, as AChRs are known to do, while others may be involved in creating the molecular scaffold for this post-synaptic specialization.

Signals produced by the nerve induce postsynaptic clusters by at least two mechanisms. First, these signals can induce redistribution of pre-existing

molecules that are initially expressed throughout the myofiber, and second, they can induce localized transcription of specific genes only by subsynaptic nuclei underlying the NMJ. Between the nerve terminal and the motor endplate is a narrow synaptic cleft containing a complex basal lamina. This basal lamina is distinguished from the adjacent extracellular matrix by the accumulation of a number of proteins, such as acetylcholinesterase and s-laminin. The synaptic basal lamina also serves as a reservoir for signaling molecules exchanged between nerve and muscle.

While the reciprocal interactions between nerve and muscle have been intensively explored for decades, many questions still remain concerning the precise nature of the signals involved in formation of the NMJ. The realization that empty sheaths of the synaptic basal lamina could induce formation of both nerve terminal specializations and motor endplates suggested that key signaling molecules might be embedded in the extracellular matrix (Sanes, J.R. et al., 1978, J.Cell. Biol. 78: 176-198; Burden, S.J., et al., 1979, J.Cell. Biol. 82: 412-425; McMahan, U.J. and Slater, C.R., 1984, J.Cell. Biol. 98: 1453-1473; Kuffler, D.P., 1986, J.Comp. Neurol. 250: 228-235). Indeed, recent findings indicate that a protein discovered for its AChR-inducing activity and thus termed ARIA (Jessell, T.M., et al., 1979, PNAS (USA) 76: 5397-5401; Falls, D.L., et al., 1990, Cold Spring Harbor Symp. Quant. Biol. 55: 397-406; Falls, D.L., et al., 1993, Cell 72: 801-815) which can increase the expression of several of the AChR subunit genes (Harris, D.A., 1989, et al., Nature 337: 173-176; Martinou, J.-C., et al., 1991, PNAS (USA) 88: 7669-7673; Jo, S.A., et al., 1995, Nature 373: 158-161; Chu, G.C., et al., 1995, Neuron 14: 329-339), is localized to the synaptic basal lamina (Jo, S.A., et al., 1995, Nature 373: 158-161; Goodearl, A.D., et al., 1995, J.Cell. Biol. 130: 1423-1434). Molecular cloning has revealed that ARIA corresponds to a factor alternatively referred to as neuregulin, NDF, heregulin or glia growth factor, and binds to the erbB family of RTKs (Carraway, K.L. and Burden, S.J., 1995, Curr. Opin. Neurobiol. 5: 606-612). Interestingly, neuregulin production has been demonstrated in

motor neurons and neuregulin receptors, erbB3 and erbB4, have recently been localized to the motor endplate, supporting the idea that nerve-derived neuregulin provides an important signal to muscle that regulates transcription from subsynaptic nuclei (Altioik, N., et al., 1995, EMBO J. 14: 4258-4266; Moscoso, L.M., et al., 1995, Dev. Biol. 172: 158-169; Zhu, X., et al., 1995, EMBO J. 14: 5842-5848).

Another protein, known as agrin, was isolated from the synaptic basal lamina based on its ability to cause redistribution of pre-existing AChRs into clusters on the surface of cultured myotubes (Godfrey, E.W., et al., 1984, J.Cell. Biol. 99: 615-627; Rupp, F., et al., 1991, Neuron 6: 811-823; Tsim, K.W., et al., 1992, Neuron 8: 677-689). In contrast to neuregulin, agrin does not appear to regulate AChR expression. However, agrin causes the clustering of a number of synaptic components, along with AChRs, in cultured myotubes (Wallace, B.G., 1989, J.Neurosci. 9: 1294-1302).

A variety of data are consistent with the notion that agrin also acts in vivo to induce and maintain the postsynaptic membrane specialization. Most important among these are the findings that the most active forms of agrin are exclusively made by neurons and are deposited in the synaptic basal lamina (Ruegg, M.A., et al., 1992, Neuron 8: 691-699; Ferns, M., et al., 1993, Neuron 11: 491-502; Hoch, W., et al., 1993, Neuron 11: 479-490), and that antibodies to agrin block nerve-induced clustering of AChRs on cultured myotubes (Reist, N.E., et al., 1992, Neuron 8: 865-868).

The precise mechanism of action of agrin remains a mystery (Sealock, R. and Froehner, S.C., 1994, Cell 77: 617-619). Agrin is known to induce tyrosine phosphorylation of AChRs, and inhibitors of tyrosine phosphorylation block agrin-mediated clustering (Wallace, B.G., et al., 1991, Neuron 6: 869-878; Wallace, B.G., 1994, J.Cell. Biol. 125: 661-668; Qu, Z. and Huganir, R.L., 1994, J.Neurosci. 14: 6834-6841; Wallace, B.G., 1995, J.Cell. Biol. 128: 1121-1129).

Intriguing recent findings have revealed that agrin can directly bind to α -dystroglycan, an extrinsic peripheral membrane protein that is attached to the cell surface by covalent linkage to β -dystroglycan, which in turn couples to the intracellular cytoskeletal scaffold via an associated protein complex (Bowe, M.A., et al., 1994, Neuron 12: 1173-1180; Campanelli, J.T., et al., 1994, Cell 77: 673-674; Gee, S.H., et al., 1994, Cell 77: 675-686; Sugiyama, J., et al., 1994, Neuron 13: 103-115; Sealock, R. and Froehner, S.C., 1994, Cell 77: 617-619).

Extrasynaptically, the dystroglycan complex binds laminin on its extracellular face, and couples to the actin scaffold via a spectrin-like molecule known as dystrophin. At the synapse however, agrin (via its own laminin-like domains) may be able to substitute for laminin, whereas utrophin (a dystrophin related protein) replaces dystrophin as the link to actin (reviewed in (Bowe, M.A. and Fallon, J.R., 1995, Ann. Rev. Neurosci. 18: 443-462)). The dystroglycan complex co-clusters with AChRs in response to agrin in vitro, and components of this complex are concentrated at the endplate in vivo (reviewed in (Bowe, M.A. and Fallon, J.R., 1995, Ann. Rev. Neurosci. 18: 443-462)).

Recent evidence suggests that a 43 kD cytoplasmic protein, known as rapsyn, anchors AChRs to a sub-synaptic cytoskeleton complex, probably via interactions with the dystroglycan complex (Cartaud, J. and Changeux, J.P., 1993, Eur. J. Neurosci. 5: 191-202; Apel, E.D., et al., 1995, Neuron 15: 115-126). Gene disruption studies reveal that rapsyn is absolutely necessary for clustering of AChRs, as well as of the dystroglycan complex. However, other aspects of NMJ formation, involving presynaptic differentiation and synapse-specific transcription, are seen in mice lacking rapsyn (Gautam, M., et al., 1995, Nature 377: 232-236).

Despite the findings that agrin can bind directly to α -dystroglycan, and that AChRs and the dystroglycan complex are linked and co-cluster in response to agrin, the role of dystroglycan as an agrin receptor remains unclear (Sealock, R. and Froehner, S.C., 1994, Cell 77: 617-619; Ferns, M., et al., 1996, J. Cell Biol. 132: 937-944). It has recently been reported that a 21kD fragment of chick agrin is sufficient to induce AChR aggregation (Gesemann, M., et al., 1995, J. Cell Biol. 128: 625-636). Dystroglycan could be directly involved in activating signaling pathways that appear to be required for clustering, such as those involving tyrosine phosphorylation, by an unknown mechanism (for example, via association with a cytoplasmic tyrosine kinase).

Alternatively, dystroglycan could be involved in couplings of agrin not only to AChRs but to a novel signaling receptor. It also remains possible that dystroglycan does not play an active or required role in initiating clustering, and is merely among an assortment of post-synaptic molecules that undergo clustering. Recent evidence indicates that the agrin fragment that is active in inducing AChR aggregation does not bind to α -dystroglycan and a structural role in aggregation, rather than a signal transfer role, has been proposed for the binding of agrin to α -dystroglycan (Gesemann, M., et al., 1996, Neuron 16: 755-767).

SUMMARY OF THE INVENTION

The present invention provides for a novel tyrosine kinase, termed "MuSK" for "muscle specific kinase," that is expressed in normal and denervated muscle, as well as other tissues including heart, spleen, ovary or retina (See Valenzuela, D., et al., 1995, Neuron 15: 573-584). The novel tyrosine kinase has alternatively been referred to as "Dmk" for "denervated muscle kinase." (See PCT International Application No. PCT/US94/08039 published 1 February 1996 as WO 96/02643 entitled Denervated Muscle Kinase (DMK), A

Receptor Of The Tyrosine Kinase Superfamily). Thus, the terms MuSK and Dmk may be used interchangeably. The protein appears to be related to the Trk family of tyrosine kinases.

5 The present invention further provides for an isolated nucleic acid molecule encoding MuSK.

The present invention also provides for a protein or peptide that comprises the extracellular domain of MuSK and the nucleic acid which encodes such
10 extracellular domain. The invention further provides for vectors comprising an isolated nucleic acid molecule encoding MuSK or its extracellular domain, which can be used to express MuSK in bacteria, yeast and mammalian cells.

The present invention also provides for use of the MuSK receptor or its
15 extracellular or intracellular domain to screen for drugs that interact with or activate MuSK. Novel agents that bind to and/or activate the receptor described herein may mediate survival, proliferation and differentiation in cells naturally expressing the receptor, but also may mediate survival, proliferation or differentiation when used to treat cells engineered to express
20 the receptor.

In particular embodiments, the extracellular domain (soluble receptor) of MuSK is utilized in screens for cognate ligands and activating molecules. For example, the MuSK receptor activating molecule described herein may be
25 used in a competition assay to identify agents capable of acting as receptor agonists or antagonists by competing the agents with MuSK activating molecule for phosphorylation of the MuSK receptor. Specifically, the active portion of human agrin described herein may be used as the MuSK activating molecule in a competition assay to screen for agents capable of acting as
30 receptor agonists or antagonists.

The term "MuSK activating molecule" as used herein refers to a molecule which is capable of inducing phosphorylation of the MuSK receptor in the context of a differentiated muscle cell. One such activating molecule is agrin as described in the Examples set forth herein.

5

The present invention also provides for nucleic acid probes, capable of hybridizing with a sequence included within the nucleotide sequence encoding human MuSK or its activating molecule, useful for the detection of MuSK expressing tissue or MuSK activating molecule-expressing tissue in humans and animals. The invention further provides for antibodies capable of specifically binding MuSK or MuSK activating molecule. The antibodies may be polyclonal or monoclonal.

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The present invention also has diagnostic and therapeutic utilities. In particular embodiments of the invention, methods of detecting aberrancies in the function or expression of the receptor described herein may be used in the diagnosis of muscular or other disorders. In other embodiments, manipulation of the receptor, agonists which bind this receptor, or receptor activating molecules may be used in the treatment of neurological diseases or diseases of muscle or neuromuscular unit disorders, including, but not limited to, muscular dystrophy and muscle atrophy. In further embodiments, the extracellular domain of the receptor is utilized as a blocking agent.

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The present invention also provides for an isolated and purified polypeptide which activates MuSK receptor. In one embodiment, the polypeptide of the invention is encoded by a nucleotide sequence comprising the coding region of the active portion of human agrin contained in the vector designated as pBluescript human Agrin-1 (pBL-hAgrin1) that was deposited with the American Type Culture Collection on December 12, 1995 under ATCC Accession No. 97378. The present invention further provides for an isolated polypeptide which is functionally equivalent to this polypeptide.

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The invention further provides for an isolated and purified nucleic acid molecule comprising a nucleotide sequence encoding the active portion of human agrin, wherein the nucleotide sequence is selected from the group consisting of:

- (a) the nucleotide sequence comprising the coding region of the active portion of human agrin contained in the vector designated as pBL-hAgrin 1 (ATCC Accession No. 97378);
- (b) a nucleotide sequence that hybridizes under stringent conditions to the nucleotide sequence of (a) and which encodes the active portion of human agrin; and
- (c) a nucleotide sequence that, as a result of the degeneracy of the genetic code, differs from the nucleotide sequence of (a) or (b) and which encodes the active portion of human agrin.

The invention also provides for the above-described nucleic acid molecule which additionally contains a nucleotide sequence so that the encoded polypeptide contains the eight amino acids ELANEIPV at the position corresponding to amino acid position 1780 as shown in Figure 14.

The invention also provides for a method of promoting the growth, survival or differentiation of a MuSK receptor expressing cell comprising administering to the MuSK receptor expressing cell an effective amount of agrin or a derivative of agrin. The method may be practiced in vitro or in vivo. In one embodiment of this method, the agrin is human agrin. In another embodiment of this method, the MuSK receptor expressing cell is a cell which is normally found in the heart, spleen, ovary, retina or skeletal muscle. In another embodiment, the MuSK receptor expressing cell is a cell which has been genetically engineered to express the MuSK receptor.

The present invention also includes a method of treating a patient suffering from a muscle disease or neuromuscular disorder comprising administering

to the patient an effective amount of agrin or a derivative thereof. By way of non-limiting example, the agrin may be human agrin and the derivative may be the active portion of the human agrin molecule.

5 The present invention also includes an antibody capable of specifically binding human agrin. More specifically, the invention includes an antibody capable of specifically binding the active portion of human agrin. The antibody may be monoclonal or polyclonal. The invention further provides a method of detecting the presence of human agrin in a sample comprising:

- 10 a) reacting the sample with an antibody capable of specifically binding human agrin under conditions whereby the antibody binds to human agrin present in the sample; and
- b) detecting the bound antibody, thereby detecting the presence of human agrin in the sample.

15 The antibody used may be monoclonal or polyclonal. The sample may be biological tissue or body fluid. The biological tissue may be brain, muscle, or spinal cord. The body fluid may be cerebrospinal fluid, urine, saliva, blood, or a blood fraction such as serum or plasma.

20 The invention further provides for an isolated and purified nucleic acid molecule comprising a nucleotide sequence encoding human muscle specific kinase (MuSK) receptor, wherein the nucleotide sequence is selected from the group consisting of:

- 25 (a) the nucleotide sequence comprising the coding region of the human MuSK receptor as set forth in Figure 4;
- (b) a nucleotide sequence that hybridizes under stringent conditions to the nucleotide sequence of (a) and which encodes a human MuSK receptor; and

- (c) a nucleotide sequence that, as a result of the degeneracy of the genetic code, differs from the nucleotide sequence of (a) or (b) and which encodes a human MuSK receptor.

5

BRIEF DESCRIPTION OF THE FIGURES

FIGURE 1 - Nucleotide and deduced amino acid (single letter code) sequences of rat musk. The nucleotide sequence encoding mature MuSK begins around nucleotide 192.

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FIGURE 2 - Northern blot showing distribution of musk in the rat during early development. Lane 1: Total embryo E9; Lane 2: Total embryo E11; Lane 3: Placenta E11; Lane 4: Embryo head E12; Lane 5: Embryo body E12; Lane 6: Embryo spinal cord E12; Lane 7: Placenta E12; Lane 8: Embryo head E13; Lane 9: Embryo body E13; Lane 10: Embryo brain E17; Lane 11: Embryo brain P1; Lane 12: Embryo brain P10; Lane 13: Embryo brain P19; Lane 14: Adult brain; Lane 15: Adult muscle; Lane 16: Adult denervated muscle; where day of sperm positivity is designated as day E1, and day of birth is designated as day P1.

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FIGURE 3 - Northern blot showing distribution of musk in adult rat tissues. Lane 1: Brain; Lane 2: Olfactory bulb; Lane 3: Cortex; Lane 4: Hippocampus; Lane 5: Thalamus/hypothalamus; Lane 6: Midbrain; Lane 7: Hindbrain; Lane 8: Cerebellum; Lane 9: Spinal Cord; Lane 10: Thymus; Lane 11: Spleen; Lane 12: Liver; Lane 13: Kidney; Lane 14: Lung; Lane 15: Sciatic Nerve; Lane 16: Retina; Lane 17: Heart; Lane 18: Ovary ; Lane 19: Muscle; Lane 20: Denervated muscle.

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FIGURE 4 - Nucleotide and deduced amino acid (single letter code) sequences of human MuSK receptor.

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FIGURE 5 - Schematic representation of genomic DNA encompassing the three kinase domain exons of the mouse MuSK gene, of the targeting vector constructed, and of a mutant locus following successful targeting. The three exons of the MuSK kinase domain are indicated as black boxes, containing the indicated kinase subdomains (SD). The PGK-neo and MC1-tk cassettes are indicated as open boxes. The novel EcoRI (R) and NcoI (N) fragments generated following successful targeting are labeled. The 5' EcoRI/HpaI probe used to detect the endogenous and mutant EcoRI fragments was derived from genomic DNA not included in the targeting construct. B, BamHI; Hp, HpaI; S, SpeI (sites included within parentheses are destroyed in the cloning process).

FIGURE 6 - MuSK Knockout Mice - Southern blot of tail DNA from wild-type, heterozygous and homozygous F2 progeny showing the endogenous and mutant EcoRI fragments detected by the 5' RI/HpaI probe, as well as the endogenous NcoI fragments detected by the kinase region probe, which are absent in the homozygous mutant.

FIGURES 7A - 7D - Post-mortem histological analysis of lung demonstrating that the alveoli air sacs in the MuSK^{-/-} newborn are not expanded (Figure 7A) as they are in the lung of the control littermate (Figure 7B), indicating that mutant pups do not take a single breath. Post-mortem histological analysis of hindlimb musculature reveals that MuSK^{-/-} mice (Figure 7C) possess grossly normal muscle architecture similar to that of control mice (Figure 7D).

FIGURES 8A - 8C - Agrin induces AChR clustering in myotubes from control but not MuSK^{-/-} mice. Myotubes derived from control and MuSK^{-/-} mice were treated overnight with varying concentrations of agrin_{4,8}, stained with rhodamine-conjugated α -bungarotoxin (α -BGT) to label surface AChRs, and then either photographed at 64X magnification under rhodamine optics (Figure 8A, challenge with 100 nM agrin depicted) or subjected to AChR

cluster quantitation (Figure 8B, each point represents the mean \pm SEM of forty myotube segments). Total AChRs on the myotubes before agrin treatment was determined by binding with ^{125}I - α -BGT (Figure 8C, each bar represents the mean \pm SEM CPM bound per μg of total cell protein (control: N=6; MuSK -/-: N= 5).

FIGURES 9A - 9D - c-agrin_{4,8} and c-agrin_{0,8} specifically induce rapid tyrosine phosphorylation of MuSK receptors. C2C12 and primary rat myoblasts were differentiated into myotubes and stimulated with conditioned media from COS cells transfected with a plasmid control (Mock) or plasmids encoding the various forms of soluble agrin, with conditioned media containing neuregulin, or with purified bFGF or insulin, as labelled. Stimulations were for ten minutes using 10 nM concentrations of the various factors, except as indicated in Figures 9C and 9D. Following factor challenges, the cells were lysed and subjected to immunoprecipitations (I.P.) for either the MuSK or ErbB3 receptors as indicated, then immunoblotted for phosphotyrosine levels. Only agrins containing the eight amino acid insert at the Z position, but not other factors, could induce MuSK phosphorylation (Figure 9A). Agrin could not induce phosphorylation of another muscle receptor, ErbB3 (Figure 9B). MuSK phosphorylation occurred at low agrin concentrations (Figure 9C) and very rapidly in response to agrin (Figure 9D).

FIGURES 10A & 10B - Agrin can not detectably bind to the isolated ectodomain of MuSK. Agrin was assayed for its binding to immobilized MuSK-Fc or to an immobilized agrin-specific monoclonal antibody (mAb), each coupled to a BIAcore sensorchip surface (Figure 10A); bindings to the MuSK-Fc surface were also done in the presence 2 mM Ca^{++} or heparin (0.01 mg/ml), as indicated, while bindings to the antibody surface were also competed with excess soluble monoclonal antibody or MuSK-Fc (each at 25

μg/ml), as indicated. Reciprocally, binding of soluble MuSK-Fc or monoclonal antibody to immobilized agrin was assayed by first binding conditioned media transfected with a plasmid control (Mock) or a plasmid encoding c-agrin_{4,8} (cAg_{4,8}) to nitrocellulose, followed by detection using either the soluble MuSK-Fc or the agrin-specific monoclonal antibody, as indicated (Figure 10B); TrkB-Fc detection of nitrocellulose-immobilized BDNF served as an additional control.

FIGURE 11 - Agrin can only induce MuSK phosphorylation in the context of a differentiated myotube: evidence for a myotube-specific accessory component. Agrin-inducible phosphorylation of an introduced chick MuSK receptor was evaluated in a clone of C2C12 myoblasts stably transfected with a chick MuSK expression vector. The introduced chick MuSK is expressed regardless of whether this C2C12 clone is undifferentiated ("Undif") or differentiated into myotubes ("Dif") (middle panel), in contrast to the endogenous mouse MuSK, which is only expressed in differentiated cells (bottom panel). However, the chick MuSK can only be inducibly phosphorylated in response to agrin when it is assayed in differentiated myotubes (top panel). The chick MuSK displays the same specificity for activation by the various agrin isoforms (each at 10 nM for ten minutes) as does the endogenous mouse MuSK (compare transfected chick MuSK and endogenous mouse MuSK in upper panel).

FIGURES 12A-12C. Relevant models for the agrin/MuSK receptor complex. Figure 12A - Schematic representation depicting the step-wise assembly of the multi-component receptor complex for ciliary neurotrophic factor (CNTF); b1, gp130; b2, LIFRb. Figure 12B - Schematic depiction of the use of soluble b receptor components (Fc-tagged) to build a CNTF receptor complex attached to the cell surface via only one of its components, the non-signaling a component; surface binding of the soluble b components can be detected using antibodies recognizing the Fc tag. Figure 12C - Schematic representation of one of several possible models of the MuSK receptor complex for agrin,

depicting requirement for a myotube-associated specificity component (M.A.S.C.) and possible interactions to additional components that may be required for signaling or coupling to various effectors or substrates; these couplings may be mediated extracellularly (for example via agrin binding to the dystroglycan complex) or intracellularly (for example via interactions of SH2 domain-containing proteins to phosphorylated tyrosines on MuSK).

FIGURES 13A - 13C. Evidence for an agrin/MuSK receptor complex utilizing a myotube-specific accessory component. Figure 13A - Formation of agrin/MuSK complexes on the surface of myotubes: undifferentiated (Undiff.) or myotube-differentiated (Diff) C2C12 cells were assayed for their ability to bind either MuSK-Fc or a control receptor-Fc fusion (TrkB-Fc), in the absence or presence of various agrin isoforms (provided in conditioned media from transient COS transfections); specific binding of MuSK-Fc to the myotube surface, which is enhanced by exogenously provided agrin, is suggested to involve complexes analogous to those depicted in Figure 12B. Figure 13B - Direct binding of agrin to MuSK is demonstrated by cross-linking analysis. Radiolabelled agrin (a recombinant c-terminal fragment (or portion) of human agrin, termed hAgrin_{4,8}) at 1 nM was chemically cross-linked to the surface of myotubes. Following cross-linking, lysates were immunoprecipitated with a MuSK-specific antibody (lane 1). The cross-linking was also done in the presence of excess (150 nM) unlabelled agrin (lane 2), while the immunoprecipitation was also done in the presence of excess peptide (corresponding to that used to generate the antibody) to block the MuSK precipitation; positions of the agrin/MuSK complex, as well as of various forms of unbound monomeric and dimeric agrin (see text), are indicated. Figure 13C - Inhibition of agrin-induced AChR clustering by MuSK-Fc: agrin-induced AChR-clustering (using 10 nM c-agrin_{4,8}) was performed on C2C12 myotube cultures in the presence of varying concentrations of soluble MuSK-Fc or a control receptor-Fc fusion (Ret-Fc); the soluble MuSK-Fc specifically inhibits, presumably by forming inactive

complexes on the cell surface with agrin and the myotube-specific accessory component.

5 FIGURE 14 - Amino acid (single letter code) sequence of rat agrin indicating Y and Z sites of amino acid inserts found in splice variants.

10 FIGURE 15 - Nucleotide and amino acid (single letter code) sequences of human agrin expression construct including the signal peptide and flg tag (FLAG tag). The start of the coding region for the active C-terminal fragment (portion) of human agrin 4-8 is indicated. Also indicated are the position Y and position Z insert sites at which the 4 and 8 amino acid inserts are located. Throughout this application, references to human agrin 4,8; c-agrin 4,8; or human c-agrin 4,8 indicate the active C-terminal fragment (portion) of human agrin 4-8 as set forth in the Figure.

15 FIGURE 16 - Results of phosphorylation assay showing that the active C-terminal 50kD portion of human agrin 4,8 and the truncated delta 9 portion of human agrin can each induce phosphorylation of the MuSK receptor.

20 FIGURE 17 - Results of pharmacokinetic study comparing serum half-lives of active C-terminal 50kD portion of human agrin 4,8 (c-agrin 4,8) with active C-terminal 50kD portion of human agrin 4,8 that has been modified by covalent addition of polyethylene glycol.

25

DETAILED DESCRIPTION OF THE INVENTION

30 The present invention provides for a novel tyrosine kinase molecule that is related to the trk family of tyrosine kinases. The sequence of the protein is set forth in Figure 1 as SEQ. ID NO: 1. The coding region of the mature protein is

believed to begin on or around the serine-glycine-threonine on or around position 20 of the coded region.

5 The novel tyrosine kinase described herein has been found to be induced in denervated skeletal muscle. Accordingly, it has been designated as MuSK (muscle specific kinase). It has also been referred to previously as Dmk (denervated muscle kinase). In addition to being found in skeletal muscle, both normal and denervated, MuSK has also been found to be present in, but not be limited to, the spleen, ovary and retina. It appears to be present during
10 early development, but is also found in adult tissue.

MuSK may be related to the Torpedo RTK identified by Jennings, et al. supra. However, MuSK differs in that it appears to be induced in denervated muscle, whereas no such induction has been reported with regard to the Torpedo
15 RTK. Furthermore, the Torpedo RTK has an extracellular kringle domain, whereas MuSK does not. However, these kinases may be members of the same or related families.

The gene encoding rat MuSK has been cloned and the DNA sequence
20 determined (Figure 1; SEQ ID NO: 2). The extracellular domain of the mature protein is believed to be encoded by the nucleotide sequence beginning on or around position 192 and ending on or around position 1610. The transmembrane portion of the protein is believed to be encoded by the nucleotide sequence beginning on or around position 1611 and ending on or
25 around position 1697. The intracellular domain is believed to be encoded by the nucleotide sequence beginning on or around position 1698 and ending on or around position 2738. A cDNA clone encoding Dmk (MuSK) was deposited with the American Type Culture Collection on July 13, 1993 and accorded an accession number of ATCC No. 75498.

30

The present invention also provides for a protein or peptide that comprises the extracellular domain of MuSK as well as the sequence of nucleotides which encode this extracellular domain. The extracellular domain of the protein is believed to be comprised of the amino acids at or around positions
5 20 through 492 of the coding region set forth as SEQ ID NO: 1.

The similarity between MuSK and the Torpedo RTK suggests the utilization of regions of sequence homologies within these genes to develop primers useful for searching for additional, related RTKs.

10 Accordingly, the invention provides for nucleic acids, or oligonucleotides greater than about 10 bases in length, that hybridize to the nucleic acid sequences described herein and that remain stably bound under stringent conditions. Stringent conditions as used herein are those which (1) employ
15 low ionic strength and high temperature for washing, for example, 0.15 M NaCl/ 0.015 M sodium citrate /0.1% NaDodSO₄ at 50°C, or (2) use during hybridization of a denaturing agent such as formamide, for example, 50% (vol/vol) formamide with 0.1% bovine serum albumin/0.1% Ficoll/0.1% polyvinylpyrrolidone/50 mM sodium phosphate buffer at pH 6.5 with 750
20 mM NaCl, 75 mM sodium citrate at 42°C.

The present invention further provides for an isolated and purified nucleic acid molecule comprising a nucleotide sequence encoding human muscle specific kinase (MuSK) receptor, wherein the nucleotide sequence is selected
25 from the group consisting of:

- (a) the nucleotide sequence comprising the coding region of the human MuSK receptor as set forth in Figure 4;
- (b) a nucleotide sequence that hybridizes under stringent conditions to the nucleotide sequence of (a) and which encodes a human MuSK
30 receptor; and

- (c) a nucleotide sequence that, as a result of the degeneracy of the genetic code, differs from the nucleotide sequence of (a) or (b) and which encodes a human MuSK receptor.

5 The invention further provides for isolated and purified human MuSK receptor encoded by the coding region of the human MuSK receptor nucleotide sequence as set forth above. The invention also provides for a vector which comprises the isolated nucleic acid molecule described. In one embodiment, the vector is an expression vector wherein the DNA molecule
10 is operatively linked to an expression control sequence. In a further embodiment, the expression vector comprises an immediate early gene promoter. In a still further embodiment, the expression vector of the invention comprises the fos promoter or the jun promoter as the early gene promoter.

15 The invention further contemplates a host-vector system for the production of a polypeptide having the biological activity of a human MuSK receptor which comprises the vector described above in a suitable host cell. By way of nonlimiting example, a suitable host cell may be a C2C12 cell or an NIH3T3
20 cell. The invention further provides for a method of producing a polypeptide having the biological activity of human MuSK receptor which comprises growing cells of the above-described host-vector system under conditions permitting production of the polypeptide and recovering the polypeptide so produced.

25 In addition, the invention provides for a therapeutic composition comprising the MuSK receptor activating molecule in a pharmaceutically acceptable vehicle.

The invention also provides for an antibody which specifically binds the above-described MuSK receptor. The antibody of the invention may be a polyclonal or monoclonal antibody.

5 The invention further provides for a MuSK receptorbody comprising the extracellular portion of the above-described MuSK receptor, fused to an immunoglobulin constant region. In a preferred embodiment, the constant region of the receptorbody is the human immunoglobulin gamma-1 constant region (MuSK-IgG1 receptorbody).

10

The invention further provides a method of detecting the presence of MuSK ligand in a sample comprising:

- a) reacting the sample with a MuSK receptorbody capable of specifically binding MuSK ligand under conditions whereby the MuSK
- 15 receptorbody binds to MuSK ligand present in the sample; and
- b) detecting the bound MuSK receptorbody, thereby detecting the presence of MuSK ligand in the sample.

The MuSK receptorbody used is most preferably MuSK-IgG1 receptorbody.

20 The sample may be biological tissue or body fluid. The biological tissue may be muscle, heart, spleen or ovary. The body fluid may be cerebrospinal fluid, urine, saliva, blood, or a blood fraction such as serum or plasma.

The invention also provides for a fibroblast cell line that is growth factor
25 dependent in serum-free medium and that comprises a nucleic acid molecule encoding the human MuSK receptor as described above.

When using nucleotide sequences coding for part or all of MuSK in
accordance with this invention to isolate new family members or MuSK from
30 other species, the length of the sequence should be at least sufficient to be capable of hybridizing with endogenous mRNA from the vertebrate's own

musk. Typically, sufficient sequence size will be about 15 consecutive bases (DNA or RNA).

Strategies for identifying novel RTKs using degenerate
 5 oligodeoxyribonucleotide primers corresponding to protein regions
 surrounding amino acids conserved in tyrosine kinases have been previously
 described (Wilks, et al., 1989, Proc. Natl. Acad. Sci. U.S.A., 86:1603-1607,
 Partanen, J. et al., 1990, Proc. Natl. Acad. Sci. U.S.A. 87: 8913-8917; Lai and
 Lemke, 1991, Neuron 6: 691-704; Masiakowski and Carroll, 1992, J. Biol. Chem.
 10 267: 26181-26190). The discovery by applicants of the relationship between
 MuSK and the Torpedo RTK has led to the identification of heretofore
 unknown homology regions which may be used in screening strategies.

The following primer, based on the amino acid homology domain Asp-Val-
 15 Trp-Ala-Tyr-Gly (SEQ ID NO: 3) between MuSK and the Torpedo RTK, may be
 used in combination with additional primers that correspond to known
 homology regions characteristic of RTKs, to isolate related tyrosine kinases,
 e.g. other family members [all codes used herein representing amino acids
 and nucleotides are as set forth in 37 C.F.R. §1.822(b)] :

20 5'-GAATTCGAGCTCCCRWANGCCCANACRTC-3' (SEQ ID NO:4)

The additional primers that correspond to known homology regions
 characteristic of RTKs include the following:

- 25 5'
- 1) Asp-Leu-Ala-Thr-Arg-Asn (SEQ ID NO: 5)
 5'-TCTTGACTCGAGAYYTNGCNACNMGNAA-3' (SEQ ID NO: 6)
 - 30 2) Asp-Leu-Ala-Ala-Arg-Asn (SEQ ID NO: 7)
 5'-TCTTGACTCGAGAYYTNGCNGCNMGNA-3' (SEQ ID NO: 8)

3'

1) Asp-Val-Trp-Ser-Leu-Gly (SEQ ID NO: 9)

3'-CTRCANACCWSNATRCCCTCGAGCTTAAG-5' (SEQ ID NO: 10)

5

2) Asp-Val-Trp-Ser-Phe-Gly (SEQ ID NO: 11)

3'-CTRCANACCWSNAARCCCTCGAGCTTAAG-5' (SEQ ID NO: 12)

3) Asp-Val-Trp-Ser-Tyr-Gly (SEQ ID NO: 13)

10

3'-CTRCANACCWSNRANCCCTCGAGCTTAAG-5' (SEQ ID NO: 14)

Alternatively, regions of homology shared by MuSK and members of related families, such as the Trk family, may be used in strategies designed to isolate novel RTKs.

15

The present invention further provides for substantially purified protein molecules comprising the amino acid sequence substantially as set forth in Figure 1 for MuSK (SEQ ID NO: 1) or functionally equivalent molecules. Functionally equivalent molecules include derivatives in which amino acid residues are substituted for residues within the sequence resulting in a silent change. For example, one or more amino acid residues within the sequence can be substituted by another amino acid of a similar polarity which acts as a functional equivalent, resulting in a silent alteration. Substitutes for an amino acid within the sequence may be selected from other members of the class to which the amino acid belongs. For example, the nonpolar (hydrophobic) amino acids include alanine, leucine, isoleucine, valine, proline, phenylalanine, tryptophan and methionine. The polar neutral amino acids include glycine, serine, threonine, cysteine, tyrosine, asparagine, and glutamine. The positively charged (basic) amino acids include arginine, lysine and histidine. The negatively charged (acidic) amino acids include aspartic acid and glutamic acid. Also included within the scope of the

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invention are proteins or fragments (portions) or derivatives thereof which are differentially modified during or after translation, e.g., by glycosylation, proteolytic cleavage, linkage to an antibody molecule or other cellular ligand, etc.

5

The invention further contemplates the isolation of proteins that have substantial similarity to the MuSK protein described herein. Substantial similarity, as used herein, refers to proteins that are from different species or are family members within a species and are identical in at least 40% of positions. Substantial similarity at the protein level includes the ability of a subject protein to compete with MuSK for binding to monoclonal antibodies raised against MuSK epitopes.

10

The MuSK protein described herein is useful in 1) screening strategies, 2) purification strategies and 3) diagnostic uses. With respect to screening strategies, expression cloning strategies based on cell survival and proliferation assays provide a method of screening for cognate ligands (Glass, et al. (1991) Cell 66:405-413). Since ligands that bind MuSK may be membrane bound, other strategies for identification of such receptors may be more well suited (Armitage, et al. 1992, Nature 357:80-82; Smith, et al. 1993, Cell 73:1349-1360). In preferred embodiments, the extracellular domain of MuSK is fused to a marker to create a chimeric protein which enables identification and purification of the extracellular domain when bound to a cognate.

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If, for example, the cognate ligand is membrane bound, as described in Smith, et al. supra, the extracellular portion of MuSK may be fused to truncated immunoglobulin heavy chains (Fc). The fusion product may then be used to identify cells expressing surface ligand that binds the receptor by, for example, flow cytometry. Alternatively, other tags, such as myc used to tag the extracellular domain of MuSK, may also be useful for the screening and

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purification of MuSK-binding ligands (Davis, et al. 1991, Science 253:59-63; Squinto, et al., 1990, Neuron 5:757-766).

5 In other embodiments, the extracellular portion of RTKs that bind known ligands are replaced with the extracellular portion of MuSK. Measurable effects, such as changes in phenotype or induction of early response genes, normally associated with binding of the known ligand to the receptor, can be used to screen for cognate ligands that induce comparable effects.

10 For example, a cell line bearing the introduced MuSK receptor or a chimeric protein comprising the extracellular domain of MuSK fused to the transmembrane domain and intracellular domain of another RTK (MuSK-chimeric receptor), as well as the parental cell line without the receptor can be exposed to any potential source of an agent that might work through the
15 receptor. Any specific effects (e.g. on cell survival or proliferation) on the cell line bearing the receptor or chimera can be used to identify and eventually purify agents acting on that receptor. Once a particular receptor/ligand system is defined, a variety of additional specific assay systems can be utilized, for example, to search for additional agonists or antagonists of MuSK.

20 According to the invention, MuSK or a MuSK-RTK chimeric receptor, when introduced into cells that do not normally express this receptor, can be used to identify ligands that bind the receptor based on the distinguishable response of the cell. The present invention contemplates that the type of response
25 elicited depends on the cell utilized, and not the specific receptor introduced into the cell. Thus, for example, expression of the MuSK receptor in PC12 pheochromocytoma cells may result in the differentiation of the PC12 cells upon exposure to a ligand that binds the receptor, whereas the same receptor in fibroblasts may mediate both survival and proliferation in response to a
30 MuSK binding ligand. Appropriate cell lines can be chosen to yield a response of the greatest utility for the assay, as well as discovery of agents that can act

on tyrosine kinase receptors. "Agents" refers to any molecule(s), including but not limited to peptide and non-peptide molecules, that will act in systems to be described in a receptor dependent manner.

5 One of the more useful systems to be exploited involves the introduction of the desired receptor into a growth factor dependent fibroblast cell line. Such a receptor which does not normally mediate proliferative responses may, following introduction into fibroblasts, nonetheless be assayed by a variety of well established methods used to quantitate effects of fibroblast growth factors
10 (e.g. thymidine incorporation or other types of proliferation assays; see van Zoelen, 1990, "The Use of Biological Assays For Detection Of Polypeptide Growth Factors" in Progress in Factor Research, Vol. 2, pp. 131-152; Zhan and M. Goldfarb, 1986, Mol. Cell. Biol., Vol. 6, pp. 3541-3544). These assays have the added advantage that any preparation can be assayed both on the cell line
15 having the introduced receptor as well as the parental cell line lacking the receptor. Only specific effects on the cell line with the receptor would be judged as being mediated through the introduced receptor.

A cell that expresses a receptor described herein may either naturally express
20 the receptor or be genetically engineered to do so. For example, nucleotide sequences obtained as described herein may be introduced into a cell by transfection, transduction, microinjection, electroporation, via a transgenic animal, etc., using any method known in the art.

25 The specific binding of test agent to the receptor may be measured in a number of ways. For example, the binding of test agent to cells may be detected or measured, by detecting or measuring (i) test agent bound to the surface of intact cells; (ii) test agent cross-linked to receptor protein in cell lysates; or (iii) test agent bound to receptor in vitro. The specific interaction
30 between test agent and the receptor may be evaluated by using reagents that demonstrate the unique properties of that interaction.

Alternatively, the specific activity of test agent on the receptor may be measured by evaluating the secondary biological effects of that activity, including, but not limited to, the induction of neurite sprouting, immediate
5 early gene expression or phosphorylation of the receptor. For example, the ability of the test agent to induce neurite sprouting can be tested in cells that lack the receptor and in comparable cells that express, for example, a chimeric receptor comprising the MuSK extracellular domain and the intracellular domain of a member of the Trk family (such as TrkA, TrkB or TrkC); neurite
10 sprouting in receptor-expressing cells but not in comparable cells that lack the receptor would be indicative of a specific test agent/receptor interaction. A similar analysis could be performed by detecting immediate early gene (e.g. fos and jun) induction in receptor-minus and receptor-plus cells, or by detecting phosphorylation of the receptor protein using standard phosphorylation
15 assays known in the art.

Similarly, the present invention provides for a method of identifying an agent that has signal transducing activity comprising (i) exposing a cell that expresses a tyrosine kinase receptor as described herein to a test agent and (ii)
20 detecting the activity of the test agent to the receptor, in which activity positively correlates with signal transducing activity. Activity may be detected by either assaying for direct binding or the secondary biological effects of binding, as discussed supra. Such a method may be particularly useful in identifying new neurotrophic factors or factors having other pharmaceutical
25 activity such as cardioprotective activity, or may be useful in screening a large array of peptide and non-peptide agents (e.g., peptidomimetics) for such activities.

In a preferred, specific, nonlimiting embodiment of the invention, a large grid
30 of culture wells may be prepared that contain, in alternate rows, PC12 (or fibroblasts, see infra) cells that are either receptor-minus or engineered to be

receptor-plus. A variety of test agents may then be added such that each column of the grid, or a portion thereof, contains a different test agent. Each well could then be scored for the presence or absence of neurite sprouting. An extremely large number of test agents could be screened for signal transducing activity in this manner.

The present invention also provides for assay systems that may be used according to the methods described supra. Such assay systems may comprise in vitro preparations of receptor, e.g. affixed to a solid support, or may preferably comprise cells that express receptor proteins described herein.

The present invention further provides for host cells and microorganisms and vectors that carry the recombinant nucleic acid molecules described supra. Cells that express receptor protein may be genetically engineered to produce receptor as described supra, by transfection, transduction, electroporation, or microinjection of nucleic acid encoding MuSK in a suitable expression vector. In one embodiment, the host cell carrying the recombinant nucleic acid is an animal cell, such as COS. In another embodiment, the host cell is a bacterium, preferably Escherichia coli.

Any of the methods known to one skilled in the art for the insertion of DNA fragments into a vector may be used to construct expression vectors encoding receptor. These methods may include in vitro recombinant DNA and synthetic techniques and in vivo recombinations (genetic recombination). Expression of nucleotide sequence encoding the receptor protein or peptide fragment may be regulated by a second nucleotide sequence so that the receptor protein or peptide is expressed in a host transformed with the recombinant DNA molecule. For example, expression of receptor may be controlled by any promoter/enhancer element known in the art. Promoters which can be used to control receptor expression include, but are not limited to the long terminal repeat as described in Squinto et al., (1991, Cell 65:1-20);

the SV40 early promoter region (Bernoist and Chambon, 1981, *Nature* 290:304-310), the CMV promoter, the M-MuLV 5' terminal repeat the promoter contained in the 3' long terminal repeat of Rous sarcoma virus (Yamamoto, et al., 1980, *Cell* 22:787-797), the herpes thymidine kinase promoter (Wagner et al., 1981, *Proc. Natl. Acad. Sci. U.S.A.* 78:144-1445), the regulatory sequences of the metallothionein gene (Brinster et al., 1982, *Nature* 296:39-42); prokaryotic expression vectors such as the β -lactamase promoter (Villa-Kamaroff, et al., 1978, *Proc. Natl. Acad. Sci. U.S.A.* 75:3727-3731), or the *tac* promoter (DeBoer, et al., 1983, *Proc. Natl. Acad. Sci. U.S.A.* 80:21-25). See also "Useful proteins from recombinant bacteria" in *Scientific American*, 1980, 242:74-94; promoter elements from yeast or other fungi such as the Gal 4 promoter, the ADH (alcohol dehydrogenase) promoter, PGK (phosphoglycerol kinase) promoter, alkaline phosphatase promoter, and the following animal transcriptional control regions, which exhibit tissue specificity and have been utilized in transgenic animals: elastase I gene control region which is active in pancreatic acinar cells (Swift et al., 1984, *Cell* 38:639-646; Ornitz et al., 1986, *Cold Spring Harbor Symp. Quant. Biol.* 50:399-409; MacDonald, 1987, *Hepatology* 7:425-515); insulin gene control region which is active in pancreatic beta cells (Hanahan, 1985, *Nature* 315:115-122), immunoglobulin gene control region which is active in lymphoid cells (Grosschedl et al., 1984, *Cell* 38:647-658; Adames et al., 1985, *Nature* 318:533-538; Alexander et al., 1987, *Mol. Cell. Biol.* 7:1436-1444), mouse mammary tumor virus control region which is active in testicular, breast, lymphoid and mast cells (Leder et al., 1986, *Cell* 45:485-495), albumin gene control region which is active in liver (Pinkert et al., 1987, *Genes and Devel.* 1:268-276), alpha-fetoprotein gene control region which is active in liver (Krumlauf et al., 1985, *Mol. Cell. Biol.* 5:1639-1648; Hammer et al., 1987, *Science* 235:53-58); alpha 1-antitrypsin gene control region which is active in the liver (Kelsey et al., 1987, *Genes and Devel.* 1:161-171), beta-globin gene control region which is active in myeloid cells (Mogam et al., 1985, *Nature* 315:338-340; Kollias et al., 1986, *Cell* 46:89-94); myelin basic protein gene control region which is active in oligodendrocyte cells in the brain (Readhead

et al., 1987, Cell 48:703-712); myosin light chain-2 gene control region which is active in skeletal muscle (Sani, 1985, Nature 314:283-286), and gonadotropic releasing hormone gene control region which is active in the hypothalamus (Mason et al., 1986, Science 234:1372-1378).

5

Expression vectors containing receptor-encoding gene inserts can be identified by three general approaches: (a) DNA-DNA hybridization, (b) presence or absence of "marker" gene functions, and (c) expression of inserted sequences. In the first approach, the presence of a foreign gene inserted in an expression
10 vector can be detected by DNA-DNA hybridization using probes comprising sequences that are homologous to an inserted gene. In the second approach, the recombinant vector/host system can be identified and selected based upon the presence or absence of certain "marker" gene functions (e.g., thymidine kinase activity, resistance to antibiotics, transformation phenotype, occlusion
15 body formation in baculovirus, etc.) caused by the insertion of foreign genes in the vector. For example, if the receptor-encoding gene is inserted within the marker gene sequence of the vector, recombinants containing the gene insert can be identified by the absence of the marker gene function. In the third approach, recombinant expression vectors can be identified by assaying
20 the foreign gene product expressed by the recombinant vector. Such assays can be based, for example, on the physical or functional properties of the receptor-encoding gene product, for example, by binding of the receptor to neurotrophic factor or to an antibody which directly recognizes the receptor. Cells of the present invention may transiently or, preferably, constitutively
25 and permanently express receptors or portions thereof.

In preferred embodiments, the present invention provides for cells that express receptors described herein or portions thereof and that also contain recombinant nucleic acid comprising an immediate early gene promoter [e.g.
30 the fos or jun promoters (Gilman et al., 1986, Mol. Cell. Biol. 6:4305-4316)]. When such a cell is exposed to a ligand that binds to the receptor, the binding

secondarily induces transcription off the immediate early promoter. Such a cell may be used to detect receptor/ligand binding by measuring the transcriptional activity of the immediate early gene promoter, for example, by nuclear run-off analysis, Northern blot analysis, or by measuring levels of a gene controlled by the promoter. The immediate early promoter may be used to control the expression of fos or jun or any detectable gene product, including, but not limited to, any of the known reporter genes, such as a gene that confers hygromycin resistance (Murphy and Efstratiadis, 1987, Proc. Natl. Acad. Sci. U.S.A. 84:8277-8281) chloramphenicol acetyltransferase (CAT), neomycin phosphotransferase (neo), beta-galactosidase beta-glucuronidase, beta-galactosidase, etc. of detecting or measuring neurotrophin activity.

Furthermore, the cells used in the assay systems of the invention may or may not be cells of the nervous system. For example, in a specific, nonlimiting embodiment of the invention, growth-factor dependent fibroblasts may be used as the basis for a signal transducing assay system. A fibroblast cell line that is growth factor dependent in serum-free media (e.g. as described in Zham and Goldfarb, 1986, Mol. Cell. Biol. 6:3541-3544) may be transfected with a receptor-encoding gene, for instance by using a CaPO₄ transfection protocol with 5 micrograms of DNA of CMV-promoter-based expression vector comprising the musk gene and one microgram of hygromycin-resistance gene-containing expression vector. After about 48 hours, the cells may then be selected for hygromycin resistance to identify positive transfectants. The cells may then be cultured for about three weeks in the presence of hygromycin, and then resistant colonies may be pooled. These cells may then be plated on tissue culture plates coated with poly-D-lysine and human fibronectin, and allowed to grow in DMEM plus 10% bovine calf serum for about four hours to allow the cells to bind to the plates. The serum-containing media may then be aspirated and the cells may be washed about three times with PBS to remove any residual serum. The cells may then be taken up with either serum free defined media (a 3:1 mixture of DMEM and

Hams F12, supplemented with 8 mM sodium bicarbonate, 15 mM HEPES, 4 x 10⁻⁶M MnCl₂, 3 mM histidine, 10⁻⁵M ethanolamine, 10⁻⁷M sodium selenite, 5 mg transferrin per liter, 200 mg bovine serum albumin-linoleic acid complex per liter gentamicin, penicillin, and streptomycin, 20 mM L-glutamine). Cells produced in this manner, then incubated with a factor capable of binding to MuSK may, after about 5 days in culture (replacing media and growth factors every 48 hours), be expected to be growing and proliferating; cells treated with an unrelated ligand at 100 ng/ml or in serum free-medium should not, however, proliferate.

Further insight into the physiological role of MuSK will come from the further definition of the activating molecule of the present invention. The kinase domain of the MuSK receptor appears to be related to other receptor tyrosine kinases, thus it is likely that the MuSK receptor is involved in signal transduction in cells in which it is expressed. Accordingly, the MuSK activating molecule of the present invention may be used to induce signal transduction not only in naturally occurring MuSK-expressing cells, which include cells found in the muscle tissue, heart, spleen, ovaries and retina, but also in cells engineered to express the MuSK receptor. The MuSK activating molecule of the present invention may be used to promote the growth or survival of such cells.

The term "MuSK activating molecule" as used herein refers to a molecule which is capable of inducing phosphorylation of the MuSK receptor in the context of a differentiated muscle cell. One such activating molecule is agrin as described in the Examples set forth herein.

As used herein, the term "MuSK activating molecule" includes the isolated and purified MuSK receptor activating polypeptides described herein, as well as functionally equivalent molecules in which amino acid residues are substituted for residues within the sequence resulting in a silent change. For

example, one or more amino acid residues within the sequence can be substituted by another amino acid of a similar polarity which acts as a functional equivalent, resulting in a silent alteration. Substitutes for an amino acid within the sequence may be selected from other members of the class to which the amino acid belongs. For example, the nonpolar (hydrophobic) amino acids include alanine, leucine, isoleucine, valine, proline, phenylalanine, tryptophan and methionine. The polar neutral amino acids include glycine, serine, threonine, cysteine, tyrosine, asparagine, and glutamine. The positively charged (basic) amino acids include arginine, lysine and histidine. The negatively charged (acidic) amino acids include aspartic acid and glutamic acid. Also included within the scope of the invention are proteins or fragments (portions) or derivatives thereof which exhibit the same or similar biological activity and derivatives which are differentially modified during or after translation, e.g., by glycosylation, proteolytic cleavage, linkage to an antibody molecule or other cellular ligand, etc.

The present invention also provides for use of the MuSK receptor or its extracellular or intracellular domain to screen for drugs that interact with or activate MuSK. Novel agents that bind to and/or activate the receptor described herein may mediate survival, proliferation and differentiation in cells naturally expressing the receptor, but also may mediate survival, proliferation or differentiation when used to treat cells engineered to express the receptor.

In particular embodiments, the extracellular domain (soluble receptor) of MuSK is utilized in screens for cognate ligands and activating molecules. For example, the MuSK receptor activating molecule described herein may be used in a competition assay to identify agents capable of acting as receptor agonists or antagonists by competing the agents with MuSK activating molecule for phosphorylation of the MuSK receptor. Specifically, the active

portion of human agrin described herein may be used as the MuSK activating molecule in a competition assay to screen for agents capable of acting as receptor agonists or antagonists.

5 The present invention also provides for nucleic acid probes, capable of hybridizing with a sequence included within the nucleotide sequence encoding human MuSK or its activating molecule, useful for the detection of MuSK expressing tissue or MuSK activating molecule-expressing tissue in humans and animals. The invention further provides for antibodies capable
10 of specifically binding MuSK or MuSK activating molecule. The antibodies may be polyclonal or monoclonal.

The present invention also has diagnostic and therapeutic utilities. In particular embodiments of the invention, methods of detecting aberrancies in
15 the function or expression of the receptor described herein may be used in the diagnosis of muscular or other disorders. In other embodiments, manipulation of the receptor, agonists which bind this receptor, or receptor activating molecules may be used in the treatment of neurological diseases or diseases of muscle or neuromuscular unit disorders, including, but not
20 limited to, muscular dystrophy and muscle atrophy. In further embodiments, the extracellular domain of the receptor is utilized as a blocking agent.

The present invention also provides for an isolated and purified polypeptide which activates MuSK receptor. In one embodiment, the polypeptide of the
25 invention is encoded by a nucleotide sequence comprising the coding region of the active portion of human agrin contained in the vector designated as pBluescript human Agrin-1 (pBL-hAgrin1) that was deposited with the American Type Culture Collection on December 12, 1995 under ATCC Accession No. 97378. The present invention further provides for an isolated
30 polypeptide which is functionally equivalent to this polypeptide.

The invention further provides for an isolated and purified nucleic acid molecule comprising a nucleotide sequence encoding the active portion of human agrin, wherein the nucleotide sequence is selected from the group consisting of:

- 5 (a) the nucleotide sequence comprising the coding region of the active portion of human agrin contained in the vector designated as pBL-hAgrin 1 (ATCC Accession No. 97378);
- (b) a nucleotide sequence that hybridizes under stringent conditions to the nucleotide sequence of (a) and which encodes the active portion of
10 human agrin; and
- (c) a nucleotide sequence that, as a result of the degeneracy of the genetic code, differs from the nucleotide sequence of (a) or (b) and which encodes the active portion of human agrin.

The invention also provides for the above-described nucleic acid molecule
15 which additionally contains a nucleotide sequence so that the encoded polypeptide contains the eight amino acids ELANEIPV at the position corresponding to amino acid position 1780 as shown in Figure 14.

The invention further provides for an isolated and purified nucleic acid
20 molecule comprising a nucleotide sequence encoding the active portion of human agrin, wherein the nucleotide sequence is selected from the group consisting of:

- (a) the nucleotide sequence comprising the coding region of the active portion of human agrin as set forth in Figure 15;
- 25 (b) a nucleotide sequence that hybridizes under stringent conditions to the nucleotide sequence of (a) and which encodes the active portion of human agrin; and
- (c) a nucleotide sequence that, as a result of the degeneracy of the genetic code, differs from the nucleotide sequence of (a) or (b) and
30 which encodes the active portion of human agrin.

The invention further provides for an isolated nucleic acid molecule comprising a nucleotide sequence encoding the active portion of human agrin, wherein the nucleotide sequence is selected from the group consisting of:

- 5 (a) the nucleotide sequence as set forth in Figure 15;
- (b) the nucleotide sequence encoding amino acids 24 to 492 as set forth in Figure 15;
- (c) the nucleotide sequence encoding amino acids 60 to 492 as set forth in Figure 15;
- 10 (d) the nucleotide sequence encoding amino acids 76 to 492 as set forth in Figure 15;
- (e) the nucleotide sequence encoding amino acids 126 to 492 as set forth in Figure 15;
- (f) the nucleotide sequence encoding amino acids 178 to 492 as set forth in Figure 15;
- 15 (g) the nucleotide sequence encoding amino acids 222 to 492 as set forth in Figure 15;
- (h) the nucleotide sequence encoding amino acids 260 to 492 as set forth in Figure 15;
- 20 (i) the nucleotide sequence encoding amino acids 300 to 492 as set forth in Figure 15;
- (j) a nucleotide sequence that hybridizes under stringent conditions to any of the nucleotide sequences of (a) through (i) and which encodes the active portion of human agrin; and
- 25 (k) a nucleotide sequence that, as a result of the degeneracy of the genetic code, differs from any of the nucleotide sequences of (a) through (j) and which encodes the active portion of human agrin.

A further embodiment of the invention is an isolated and purified nucleic acid molecule encoding agrin 0-8 comprising a nucleotide sequence encoding the active portion of human agrin, wherein the nucleotide sequence is as set

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forth in Figure 15 with the exception that there is no insert at position Y.
Another embodiment of the invention is an isolated and purified nucleic acid molecule encoding agrin 4-0 comprising a nucleotide sequence encoding the active portion of human agrin, wherein the nucleotide sequence is as set forth
5 in Figure 15 with the exception that there is no insert at position Z.

The present invention provides for an isolated polypeptide encoded by any one of the nucleic acid molecules of the invention as set forth herein.
Furthermore, the present invention provides for said polypeptides modified
10 by covalent attachment of a polyethylene glycol molecule.

Thus, the present invention provides truncated forms of the human agrin polypeptide which retain one or more of the biological activities of human agrin. As set forth herein, the invention also provides nucleic acid sequences
15 encoding such truncated forms. These truncated forms retain, for example, the ability to induce phosphorylation of the MuSK receptor. The truncated forms may be of any suitable length, as long as they retain one or more of the biological activities of human agrin. Truncated forms including the C-terminal of human agrin are preferred.

20 Referring to Figure 15, starting at the N-terminal end (amino acid 24 - KSPC) these truncated forms of human agrin preferably have deletions of up to 10, 20, 30, 40, 50, 100, 150, 200, 250, 300, 350 or 400 amino acids. Particularly preferred truncated forms are described herein as delta 3 through delta 9.

25 The invention also provides for a method of promoting the growth or survival of a MuSK receptor expressing cell in culture comprising administering to the MuSK receptor expressing cell an effective amount of agrin or a derivative of agrin. In one embodiment of this method, the agrin is
30 human agrin. In another embodiment of this method, the MuSK receptor expressing cell is a cell which is normally found in the heart, spleen, ovary or

retina. In another embodiment, the MuSK receptor expressing cell is a cell which has been genetically engineered to express the MuSK receptor.

5 The present invention also includes a method of treating a patient suffering from a muscle disease or neuromuscular disorder comprising administering to the patient an effective amount of agrin or a derivative thereof. By way of non-limiting example, the agrin may be human agrin and the derivative may be the active portion of the human agrin molecule. The active portion of the human agrin molecule may be any one of the truncated fragments (portions)
10 of human agrin as described herein that is capable of inducing phosphorylation of the MuSK receptor.

The present invention also includes a method of treating a patient suffering from a muscle disease or neuromuscular disorder comprising administering
15 to the patient an effective amount of agrin or a portion or derivative thereof in combination with Ciliary Neurotrophic Factor (CNTF) or Modified Ciliary Neurotrophic Factor as described in United States Patent No. 5,349,056 issued September 20, 1994 to Panayotatos.

20 The present invention also includes an antibody capable of specifically binding human agrin. More specifically, the invention includes an antibody capable of specifically binding the active portion of human agrin. The antibody may be monoclonal or polyclonal. The invention further provides a method of detecting the presence of human agrin in a sample comprising:

- 25 a) reacting the sample with an antibody capable of specifically binding human agrin under conditions whereby the antibody binds to human agrin present in the sample; and
b) detecting the bound antibody, thereby detecting the presence of human agrin in the sample.

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The antibody used may be monoclonal or polyclonal. The sample may be biological tissue or body fluid. The biological tissue may be brain, muscle, or spinal cord. The body fluid may be cerebrospinal fluid, urine, saliva, blood, or a blood fraction such as serum or plasma.

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The cDNA clone encoding the active portion of human agrin described herein will facilitate screening of cDNA and genomic libraries in order to clone the full length sequence coding for the entire human agrin molecule. Cells may be genetically engineered to produce the active portion or the full length agrin molecule by, e.g., transfection, transduction, electroporation, microinjection, via a transgenic animal, of a nucleotide sequence encoding the active portion or the full length agrin molecule in a suitable expression vector. The invention also provides for a vector comprising an isolated nucleic acid molecule encoding an active portion or the full length human agrin molecule.

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The invention further provides for a host-vector system for the production in a suitable host cell of a polypeptide having the biological activity of human agrin. The suitable host cell may be a bacterial cell such as E. coli, a yeast cell such as Pichia pastoris, an insect cell such as Spodoptera frugiperda or a mammalian cell such as a COS or CHO cell. The invention also provides for a method of producing a polypeptide having the biological activity of human agrin which comprises growing cells of the host-vector system under conditions permitting production of the polypeptide and recovering the polypeptide so produced.

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The invention further provides for an expression vector comprising a nucleic acid molecule encoding human agrin or a portion thereof, wherein the nucleic acid molecule is operatively linked to an expression control sequence. The invention also provides a host-vector system for the production of a polypeptide having the biological activity of human agrin which comprises

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the expression vector of the invention in a suitable host cell. The suitable host cell may be a bacterial cell such as E. coli, a yeast cell such as Pichia pastoris, an insect cell such as Spodoptera frugiperda or a mammalian cell such as a COS or CHO cell. The invention further provides for a method of
5 producing a polypeptide having the biological activity of human agrin which comprises growing cells of the host-vector system of the invention, under conditions permitting production of the polypeptide and recovering the polypeptide so produced.

10 As described above, the present invention relates to a tyrosine kinase receptor that appears to be expressed in denervated muscle. According to the present invention, probes capable of recognizing these receptors may be used to identify diseases or disorders by measuring altered levels of the receptor in cells and tissues. Such diseases or disorders may, in turn, be treatable using
15 the activating molecule disclosed herein. Such disorders include but are not limited to those in which atrophic or dystrophic change of muscle is the fundamental pathological finding. For example, muscle atrophy can result from denervation (loss of contact by the muscle with its nerve) due to nerve trauma; degenerative, metabolic or inflammatory neuropathy (e.g. Guillian-Barre syndrome), peripheral neuropathy, or damage to nerves caused by
20 environmental toxins or drugs. In another embodiment, the muscle atrophy results from denervation due to a motor neuronopathy. Such motor neuronopathies include, but are not limited to: adult motor neuron disease, including Amyotrophic Lateral Sclerosis (ALS or Lou Gehrig's disease);
25 infantile and juvenile spinal muscular atrophies, and autoimmune motor neuropathy with multifocal conduction block. In another embodiment, the muscle atrophy results from chronic disuse. Such disuse atrophy may stem from conditions including, but not limited to: paralysis due to stroke, spinal cord injury; skeletal immobilization due to trauma (such as fracture, sprain or
30 dislocation) or prolonged bed rest. In yet another embodiment, the muscle atrophy results from metabolic stress or nutritional insufficiency, including,

but not limited to, the cachexia of cancer and other chronic illnesses, fasting or rhabdomyolysis, endocrine disorders such as, but not limited to, disorders of the thyroid gland and diabetes. The muscle atrophy can also be due to a muscular dystrophy syndrome, including but not limited to the Duchenne, Becker, myotonic, Fascioscapulohumeral, Emery-Dreifuss, oculopharyngeal, scapulohumeral, limb girdle, and congenital types, and the dystrophy known as Hereditary Distal Myopathy. In a further embodiment, the muscle atrophy is due to a congenital myopathy, including, but not limited to Benign Congenital Hypotonia, Central Core disease, Nemaline Myopathy, and Myotubular (centronuclear) myopathy. In addition, MuSK and its associated ligand may be of use in the treatment of acquired (toxic or inflammatory) myopathies. Myopathies which occur as a consequence of an inflammatory disease of muscle, include, but not limited to polymyositis and dermatomyositis. Toxic myopathies may be due to agents, including, but are not limited to adiodarone, chloroquine, clofibrate, colchicine, doxorubicin, ethanol, hydroxychloroquine, organophosphates, perihexiline, and vincristine.

Although not wishing to be bound by theory, preliminary mapping of musk in the mouse has revealed that the gene is localized to mouse chromosome 4 in a region of homology with human chromosome 9q. Mutations in mice that are associated with this region of chromosome 4 include the "wi" mutation (whirler), which results in symptoms of the shaker syndrome, including deafness, head-tossing, circling and hyperactivity (Lane, P.W., 1963, J. Hered. 54:263-266). Another mutation in mice that is associated with this region of chromosome 4 is the "vc" mutation (vacillans) which is associated with the symptoms of violent tremor when walking and with swaying of the hindquarters (Sirlin, J.L., 1956, J. Genet. 54:42-48).

In humans, the disease known as idiopathic torsion dystonia (ITD) is associated with a gene that has been mapped, through linkage analysis to

human chromosome 9q band 34. This disease is characterized by sustained, involuntary muscle contractions, frequently causing twisting and repetitive movements or abnormal postures.

5 Assuming a defect in musk to be associated with these diseases, the present invention may be used in gene therapy for the replacement of such gene in situ. Alternatively, probes utilizing a unique segment of the musk gene may prove useful as a diagnostic for such disorders. The present invention may also be used, where indicated, in gene therapy for the replacement of the
10 human agrin gene in situ.

Any of the methods known to one skilled in the art of transferring genes into skeletal muscle tissue may be used in MuSK or agrin gene therapy protocols. By way of non-limiting example, one skilled in the art may utilize direct
15 injection of naked DNA into muscle tissue, adenovirus-associated gene transfer, primary myoblast transplantation, or cationic liposome: DNA complex gene transfer.

For example, direct injection of DNA into muscle may be employed to
20 optimize vector construction as described by Manthorpe et al., (1993, Hum. Gene Ther. 4: 419-431) in which covalently closed circular plasmid DNA encoding the firefly luciferase reporter gene was injected into adult murine skeletal muscle for the purpose of evaluating the efficacy of various regulatory elements contained in the DNA expression vector. In a biological
25 study, the systemic immunological effects of cytokine genes were evaluated using direct injection into muscle of DNA encoding the genes for IL-2, IL-4, or TGF- β -1 (Raz, et al., 1993, Proc. Natl. Acad. Sci. USA 90: 4523-7). Another study tested the ability of the human kallikrein gene product to reduce blood pressure in spontaneously hypertensive rats following direct DNA injection
30 of the human kallikrein gene into murine skeletal muscle (Xiong, et al., 1995,

Hypertension 25: 715-719). In studies aimed at evaluating the expression of muscle-specific proteins following direct injection of DNA various deletion-containing dystrophin gene mutant DNA constructs were injected into mdx mouse skeletal muscle and expression patterns and colocalization studies were performed to investigate dystrophin function (Fritz, et al., 1995, *Pediatr. Res.* 37: 693-700). Many investigators believe that it may be not only important but advantageous to regulate the timing and level of gene expression following gene transfer through direct DNA injection. For example, Dhawan et al., (1995, *Somat. Cell. Mol. Genet.* 21: 233-240) have tested a tetracycline-responsive promoter system in which orally or parenterally administered tetracycline can regulate reporter gene expression in mouse skeletal muscle following direct injection of DNA.

Adenovirus-mediated in vivo gene transfer has been studied extensively as a possible method for delivering genes for gene therapy. Recombinant adenovirus vectors containing exogenous genes for transfer are derived from adenovirus type 5 and are rendered replication-deficient by deletion of the E1 region of the viral genome (Brody & Crystal, 1994, *Ann. N. Y. Acad. Sci.* 716: 90-101). Huard et al., (1995, *Gene Ther.* 2:107-115) have evaluated the efficiency of viral transduction into rat tissues following various routes of administration (intra-arterial, intravenous, gastric-rectal, intraperitoneal, and intracardiac). The investigators report that route of administration is a major determinant of the transduction efficiency of rat tissue by adenovirus recombinants. In addition to route of administration preferences, it has been shown that vectors carrying U3 region viral long terminal repeats (LTRs) modified in the enhancer region may be used to target tissue- and differentiation-specific gene expression into skeletal muscle (Ferrari et al., 1995, *Hum. Gene Ther.* 6: 733-742). Many studies have been performed (Ragot, et al., 1993, *Nature* 361: 647-50; Petrof, et al., 1995, *Am. J. Respir. Cell. Mol. Biol.* 13: 508-17; Phelps, et al., 1995, *Hum. Mol. Genet* 4: 1251-1258; Kochanek, et al., 1996, *Proc. Natl. Acad. Sci. USA* 93: 5731-36) which have

tested adenovirus-mediated transfer of full length and truncated forms of the dystrophin gene into muscle of normal and mdx mice.

Transplantation into skeletal muscle tissue of retrovirally transformed
5 primary myoblasts expressing recombinant genes has also been extensively
studied as a possible approach to gene therapy for muscle as well as non-
muscle diseases. It is known that the success of myoblast transplantation for
correction of intrinsic muscle defects is dependent on the ability of the
transplanted myoblasts to fuse to the host myofibers. To address this issue,
10 Rando & Blau (1994, J. Cell. Biol. 125:1275-87) developed a novel culture
system for isolating enriched and clonal populations of primary myoblasts.
Myoblasts isolated by this technique were shown to efficiently fuse to host
myofibers to form hybrid myofibers persisting for up to six months as
evidenced by β -galactosidase reporter gene expression. Because
15 immunorejection of transplanted myoblasts is a potential problem in this
gene therapy approach, it has been addressed in studies comparing autologous
versus heterologous myoblasts for transplantation (Huard, et al., 1994, Hum.
Gene Ther. 5: 949-58) and with Cyclosporin A-induced immunosuppression
in adult mice receiving myoblast transplants following muscle injury
20 (Irintchev et al., 1995, J. Neurocytol. 24: 319-331). Representative disease
targets for gene therapy using myoblast transplantation include hemophilia B
in which circulating human or canine factor IX has been measured in the
plasma of mice following transplantation of recombinant myoblasts into
skeletal muscles of normal and SCID mice (Roman, et al., 1992, Somat. Cell.
25 Mol. Genet. 18: 247-58; Dai, et al., 1992, Proc. Natl. Acad. Sci. USA 89: 10892-5;
Yao, et al., Proc. Natl. Acad. Sci. USA 89: 3357-61; Yao, et al., 1994, Gene Ther. 1:
99-107) and primary myopathies such as Duchenne muscular dystrophy
where myoblasts expressing the dystrophin gene have been transplanted into
normal and mdx mice (Partridge, et al., Nature 337: 176-9; Sopper, et al., 1994,
30 Gene Ther. 1: 108-113).

Plasmid DNA complexed with cationic lipids has been evaluated for its ability to deliver genes into muscle tissue as well. Trivedi, et al., (1995, J.

Neurochem. 64: 2230-38) carried out in vitro studies that utilized polycationic liposomes to successfully deliver the reporter gene LacZ into the cultured mouse myoblast cell line C2C12 and into primary mouse myoblasts derived from normal and mdx mice, forming the basis for adaptation to in vivo gene therapy.

The present invention provides for a method of diagnosing a neurological or other disorder in a patient comprising comparing the levels of expression of MuSK in a patient sample with the levels of expression of MuSK in a comparable sample from a healthy person, in which a difference in the levels of expression of MuSK in the patient compared to the healthy person

indicates that a disorder in the patient may be primarily or secondarily related to MuSK metabolism. A patient sample may be any cell, tissue, or body fluid but is preferably muscle tissue, cerebrospinal fluid, blood, or a blood fraction such as serum or plasma.

One variety of probe which may be used is anti-MuSK antibody or fragments thereof containing the binding domain of the antibody.

According to the invention, MuSK protein, or fragments or derivatives thereof, may be used as an immunogen to generate anti-MuSK antibodies. By providing for the production of relatively abundant amounts of MuSK protein using recombinant techniques for protein synthesis (based upon the MuSK nucleotide sequences of the invention), the problem of limited quantities of MuSK has been obviated.

To further improve the likelihood of producing an anti-MuSK immune response, the amino acid sequence of MuSK may be analyzed in order to

identify portions of the molecule which may be associated with increased immunogenicity. For example, the amino acid sequence may be subjected to computer analysis to identify surface epitopes which present computer-generated plots of hydrophilicity, surface probability, flexibility, antigenic index, amphiphilic helix, amphiphilic sheet, and secondary structure of MuSK. Alternatively, the deduced amino acid sequences of MuSK from different species could be compared, and relatively non-homologous regions identified; these non-homologous regions would be more likely to be immunogenic across various species.

For preparation of monoclonal antibodies directed toward MuSK, or its activating molecule, any technique which provides for the production of antibody molecules by continuous cell lines in culture may be used. For example, the hybridoma technique originally developed by Kohler and Milstein (1975, *Nature* 256:495-497), as well as the trioma technique, the human B-cell hybridoma technique (Kozbor et al., 1983, *Immunology Today* 4:72), and the EBV-hybridoma technique to produce human monoclonal antibodies (Cole et al., 1985, in "Monoclonal Antibodies and Cancer Therapy," Alan R. Liss, Inc. pp. 77-96) and the like are within the scope of the present invention.

The monoclonal antibodies for therapeutic use may be human monoclonal antibodies or chimeric human-mouse (or other species) monoclonal antibodies. Human monoclonal antibodies may be made by any of numerous techniques known in the art (e.g., Teng et al., 1983, *Proc. Natl. Acad. Sci. U.S.A.* 80:7308-7312; Kozbor et al., 1983, *Immunology Today* 4:72-79; Olsson et al., 1982, *Meth. Enzymol.* 92:3-16). Chimeric antibody molecules may be prepared containing a mouse antigen-binding domain with human constant regions (Morrison et al., 1984, *Proc. Natl. Acad. Sci. U.S.A.* 81:6851, Takeda et al., 1985, *Nature* 314:452).

Various procedures known in the art may be used for the production of polyclonal antibodies to epitopes of MuSK. For the production of antibody, various host animals can be immunized by injection with MuSK protein, or a fragment or derivative thereof, including but not limited to rabbits, mice, rats, etc. Various adjuvants may be used to increase the immunological response, depending on the host species, and including but not limited to Freund's (complete and incomplete), mineral gels such as aluminum hydroxide, surface active substances such as lysolecithin, pluronic polyols, polyanions, peptides, oil emulsions, keyhole limpet hemocyanins, dinitrophenol, and potentially useful human adjuvants such as BCG (Bacille Calmette-Guerin) and Corynebacterium parvum.

A molecular clone of an antibody to a MuSK epitope can be prepared by known techniques. Recombinant DNA methodology (see e.g., Maniatis et al., 1982, Molecular Cloning, A Laboratory Manual, Cold Spring Harbor Laboratory, Cold Spring Harbor, New York) may be used to construct nucleotide sequences which encode a monoclonal antibody molecule, or antigen binding region thereof.

Antibody molecules may be purified by known techniques, e.g., immunoabsorption or immunoaffinity chromatography, chromatographic methods such as HPLC (high performance liquid chromatography), or a combination thereof, etc.

The present invention provides for antibody molecules as well as fragments of such antibody molecules. Antibody fragments which contain the idiotype of the molecule can be generated by known techniques. For example, such fragments include but are not limited to: the F(ab')₂ fragment which can be produced by pepsin digestion of the antibody molecule; the Fab' fragments which can be generated by reducing the disulfide bridges of the F(ab')₂

fragment, and the Fab fragments which can be generated by treating the antibody molecule with papain and a reducing agent.

The above mentioned probes may be used experimentally to identify cells or tissues which hitherto had not been shown to express musk. Furthermore, these methods may be used to identify the expression of musk by aberrant tissues, such as malignancies. In additional embodiments, these methods may be used diagnostically to compare the expression of musk in cells, fluids, or tissue from a patient suffering from a disorder with comparable cells, fluid, or tissue from a healthy person. Fluid is construed to refer to any body fluid, but particularly blood, including blood fractions such as serum or plasma, or cerebrospinal fluid. A difference in the levels of expression of musk in the patient compared to a healthy person may indicate that the patient's disorder may be primarily or secondarily related to MuSK metabolism. An increase in levels of MuSK, for example, could either indicate that the patient's disorder is associated with an increased sensitivity to normal levels of MuSK-binding ligand or, alternatively, may suggest that the patient's MuSK-binding ligand levels are low such that the number of receptors is increased by way of compensation.

The present invention further provides for the use of soluble receptor (the extracellular domain) to counter the effect of ligand on MuSK expressing cells.

EXAMPLE 1 - CLONING OF THE cDNA ENCODING MuSK

Tyrosine kinase homology domains were identified based on the alignments by Hanks et al. (1988) Science 241, 42-52. Highly conserved regions Asp-Leu-Ala-Ala-Arg-Asn (SEQ ID NO: 7) AND Asp-Val-Trp-Ser-Tyr-Gly (SEQ ID NO: 13) were used in designing the following degenerate oligonucleotide primers:

5'-TCTTGACTCGAGAYYTNGCNGCNMGNA-3' (SEQ ID NO: 8)

5'-GAATTCGAGCTCCCRANSWCCANACRTC-3' (SEQ ID NO: 15)

with which to prime PCR reactions using denervated muscle cDNAs.

Resulting amplified DNA fragments were cloned by insertion into plasmids,
sequenced and the DNA sequences were compared with those of all known
tyrosine kinases. cDNA templates were generated by reverse transcription of
denervated muscle tissue RNAs using oligo d(T) primers. PCR reactions were
done at primer annealing temperatures of 40°C. Aliquots of the PCR reactions
were subjected to electrophoresis on an agarose gel.

Size-selected amplified DNA fragments from these PCR reactions were cloned
into plasmids as follows: Each PCR reaction was reamplified as described
above, digested with XhoI and SacI to cleave sites in the termini of the
primers (see below). XhoI/SacI-cut DNAs were purified by Magic PCR kit
(from Promega) and cloned into compatible XhoI/SacI sites in the Bluescript II
SK(+) plasmid, introduced into DH10B E. coli by electroporation, followed by
plating of transformants on selective agar. Ampicillin-resistant bacterial
colonies from PCR transformation were inoculated into 96-well microtiter
plates and used for PCR using vector primers (T3 and T7) flanking the
tyrosine kinase insert and these PCR fragments were analyzed by sequencing.

One of the cloned fragment sequences contained a segment of a novel
tyrosine kinase domain, which was designated as MuSK. The sequence of the
PCR-derived fragment corresponding to MuSK was used to generate PCR
primers to obtain longer MuSK specific fragments by the RACE procedure.
These longer MuSK probes were used as a hybridization probe to obtain full
length MuSK cDNA clones from a rat denervated skeletal muscle cDNA
library. DNA was sequenced by using the ABI 373A DNA sequencer and Taq
Dyedeoxy Terminator Cycle Sequencing Kit (Applied Biosystems, Inc., Foster
City, CA). The sequence of MuSK (Figure 1; SEQ ID NO:1) has a high degree of

homology to members of the trk family of proteins. It was also found to be similar to the Jennings, et al. Torpedo RTK found in muscle.

Oligonucleotide primers corresponding to conserved regions of known
5 tyrosine kinase molecules were used to amplify and clone DNA sequences
encoding novel orphan tyrosine kinase receptor molecules. The amino acid
sequences of representatives from branches of the tyrosine kinase family and
regions of homology within the catalytic domain of these proteins were used
to design degenerate oligonucleotide primers. These primers were then used
10 to prime PCR reactions using as template a rat denervated muscle cDNA
library. Resulting amplified DNA fragments were then cloned into Bluescript
II SK(+) plasmid, sequenced, and the DNA sequences compared with those of
known tyrosine kinases. The sequence of a PCR fragment encoding a novel
tyrosine kinase named MuSK was used to obtain more adjoining DNA
15 sequence. A DNA fragment containing MuSK sequences was used as a probe
to obtain a cDNA clone from a denervated skeletal muscle library. This clone
encodes a novel tyrosine kinase receptor with a high degree of homology to
members of the trk family of proteins. It was also found to be homologous to
the Jennings, et al. Torpedo RTK. Figure 1 presents the nucleotide sequence
20 (SEQ ID NO: 2) of the musk clone.

EXAMPLE 2 - IDENTIFICATION OF ADDITIONAL TYROSINE KINASES

25 The novel MuSK sequence is used to obtain homology segments among
receptor tyrosine kinases which can be used in combination with other
homology segments. For example, an alignment of the Torpedo trk-related
kinase with MuSK shows the following conserved protein segment:

Asp-Val-Trp-Ala-Tyr-Gly (SEQ ID NO: 3)

30 This homology "box" is not present in any other mammalian tyrosine kinase
receptor. Degenerated oligonucleotides essentially based on this "box" in

combination with either previously known or novel tyrosine kinase
homology segments can be used to identify new tyrosine kinase receptors.

The highly conserved regions between MuSK and Torpedo TRK Asp-Val-Trp-
Ala-Tyr-Gly (SEQ ID NO: 3) as well as additional primers based on known
regions of homology, such as SEQ ID NOS. 5, 7, 9 OR 11, are used in designing
degenerate oligonucleotide primers with which to prime PCR reactions using
cDNAs. cDNA templates are generated by reverse transcription of tissue
RNAs using oligo d(T) or other appropriate primers. Aliquots of the PCR
reactions are subjected to electrophoresis on an agarose gel. Resulting
amplified DNA fragments are cloned by insertion into plasmids, sequenced
and the DNA sequences are compared with those of all known tyrosine
kinases.

Size-selected amplified DNA fragments from these PCR reactions are cloned
into plasmids as follows. Each PCR reaction is reamplified as described above
in Example 1, digested with XhoI and SacI to cleave sites in the termini of the
primers (see below). XhoI/SacI-cut DNAs are cloned into compatible
XhoI/SacI sites in a plasmid, introduced into E. coli by electroporation,
followed by plating of transformants on selective agar. Ampicillin-resistant
bacterial colonies from PCR transformation are inoculated into 96-well
microtiter plates and individual colonies from these PCR clones are analyzed
by sequencing of plasmid DNAs that are purified by standard plasmid
miniprep procedures.

Cloned fragments containing a segment of a novel tyrosine kinase domain
are used as hybridization probes to obtain full length cDNA clones from a
cDNA library.

EXAMPLE 3 - TISSUE SPECIFIC EXPRESSION OF MuSK

A 680 nts fragment, containing the tyrosine kinase domain of MuSK, was radiolabeled and utilized in Northern analysis of various rat tissue specific RNAs. The rat tissue specific RNAs were fractionated by electrophoresis through a 1% agarose-formaldehyde gel followed by capillary transfer to a nylon membrane with 10X SSC. The RNAs were cross-linked to the membranes by exposure to ultraviolet light and hybridized at 65°C to the radiolabeled MuSK probe in the presence of 0.5M NaPO₄ (pH 7), 1% bovine serum albumin (Fraction V, Sigma), 7% SDS, 1 mM EDTA and 100 ng/ml sonicated, denatured salmon sperm DNA. The filter was washed at 65°C with 2X SSC, 0.1% SDS and subjected to autoradiography for 5 days with one intensifying screen and X-ray film at -70°C. Ethidium bromide staining of the gel demonstrated that equivalent levels of total RNA were being assayed for the different samples.

The MuSK probe hybridized strongly in adult rat tissue (Figure 3) to a 7 kb transcript from denervated skeletal muscle, and weakly to normal muscle, retina, ovary, heart and spleen. Weaker levels of expression could also be found in liver, kidney and lung. It also hybridizes weakly to a shorter MuSK transcript of about 6 kb in brain, spinal cord and cerebellum.

In embryonic tissue (Figure 2), MuSK transcripts can be found in body, spinal cord, placenta and head at E12 and E 13.

The high expression of MuSK in muscle and neural tissue suggests that the present invention may be utilized to treat disorders of the nervous system, specifically the wide array of neurological disorders affecting motor neurons (see discussion, supra) and the neuromuscular junction. Additionally, high expression of MuSK in heart tissue suggests that the present invention may be utilized to treat heart disease, and may, for example, have prophylactic use

in preventing muscle loss during or following a cardiac event. (see discussion, supra). Expression of MuSK in retinal tissue suggests that the present invention may be utilized to treat retina related disorders, including but not limited to retinitis pigmentosa. Expression of MuSK in ovaries suggests that MuSK or the ligand associated with MuSK may be useful in the treatment of diseases or disorders involving the ovaries. Finally, expression of MuSK in spleen suggests that MuSK or the ligand associated with MuSK may be useful in the treatment of diseases or disorders involving the spleen.

EXAMPLE 4 - CLONING AND EXPRESSION OF MuSK RECEPTORBODY FOR AFFINITY-BASED STUDY OF MuSK LIGAND INTERACTIONS

An expression construct was created that would yield a secreted protein consisting of the entire extracellular portion of the rat MuSK receptor fused to the human immunoglobulin gamma-1 constant region (IgG1 Fc). This fusion protein is called a Dmk or MuSK "receptorbody" (RB), and would be normally expected to exist as a dimer in solution based on formation of disulfide linkages between individual IgG1 Fc tails. The Fc portion of the MuSK RB was prepared as follows. A DNA fragment encoding the Fc portion of human IgG1 that spans from the hinge region to the carboxy-terminus of the protein, was amplified from human placental cDNA by PCR with oligonucleotides corresponding to the published sequence of human IgG1; the resulting DNA fragment was cloned in a plasmid vector. Appropriate DNA restriction fragments from a plasmid encoding MuSK receptor and from the human IgG1 Fc plasmid were ligated on either side of a short PCR-derived fragment that was designed so as to fuse, in-frame, the MuSK and human IgG1 Fc protein-coding sequences. Thus, the resulting MuSK ectodomain-Fc fusion protein precisely substituted the IgG1 Fc in place of the region spanning the

MuSK transmembrane and cytoplasmic domains. An alternative method of preparing receptorbodies is described in Goodwin, et. al. Cell 73: 447-456 (1993).

5 Milligram quantities of MuSK RB were obtained by cloning the MuSK RB
DNA fragment into the pVL1393 baculovirus vector and subsequently
infecting the Spodoptera frugiperda SF-21AE insect cell line. Alternatively,
the cell line SF-9 (ATCC Accession No. CRL-1711) or the cell line BTI-TN-5b1-
4 may be used. DNA encoding the MuSK RB was cloned as an Eco RI-NotI
10 fragment into the baculovirus transfer plasmid pVL1393. Plasmid DNA
purified by cesium chloride density gradient centrifugation was recombined
into viral DNA by mixing 3 mg of plasmid DNA with 0.5 mg of Baculo-Gold
DNA (Pharminigen), followed by introduction into liposomes using 30mg
Lipofectin (GIBCO-BRL). DNA-liposome mixtures were added to SF-21AE
15 cells (2x 10⁶ cells/60mm dish) in TMN-FH medium (Modified Grace's Insect
Cell Medium (GIBCO-BRL) for 5 hours at 27°C, followed by incubation at 27°C
for 5 days in TMN-FH medium supplemented with 5% fetal calf serum.
Tissue culture medium was harvested for plaque purification of recombinant
viruses, which was carried out using methods previously described (O'Reilly,
D.R., L.K. Miller, and V.A. Luckow, Baculovirus Expression Vectors- A
20 Laboratory Manual. 1992, New York: W.H. Freeman) except that the agarose
overlay contained 125 mg/mL X-gal (5-bromo-4-chloro-3-indolyl-b-D-
galactopyranoside; GIBCO-BRL). After 5 days of incubation at 27°C, non-
recombinant plaques were scored by positive chromagenic reaction to the X-
gal substrate, and their positions marked. Recombinant plaques were then
25 visualized by addition of a second overlay containing 100 mg/mL MTT (3-[4,5-
dimethylthiazol-2-yl]2,5-diphenyltetrazolium bromide; Sigma). Putative
recombinant virus plaques were picked by plug aspiration, and purified by
multiple rounds of plaque isolation to assure homogeneity. Virus stocks
were generated by serial, low-multiplicity passage of plaque-purified virus.
30 Low passage stocks of one virus clone (vMuSK receptor body) were produced.

SF-21AE cells were cultured in serum free medium (SF-900 II, Gibco BRL) containing 1X antibiotic/antimycotic solution (Gibco BRL) and 25 mg/L Gentamycin (Gibco BRL). Pluronic F-68 was added as a surfactant to a final concentration of 1g/L. Cultures (4L) were raised in a bioreactor (Artisan Cell Station System) for at least three days prior to infection. Cells were grown at 27°C, with gassing to 50 % dissolved oxygen, at a gas flow rate of 80 mL/min (aeration at a sparge ring). Agitation was by means of a marine impeller at a rate of 100 rpm. Cells were harvested in mid-logarithmic growth phase (~2 X 10⁶ cells per mL), concentrated by centrifugation, and infected with 5 plaque forming units of vMuSK Receptor Body per cell. Cells and inoculum were brought to 400mL with fresh medium, and virus was adsorbed for 2 hours at 27°C in a spinner flask. The culture was then resuspended in a final volume of 8L with fresh serum-free medium, and the cells incubated in the bioreactor using the previously described conditions.

Culture medium from vMuSK Receptor Body-infected SF21AE cells were collected by centrifugation (500x g, 10 minutes) at 72 hours post-infection. Cell supernatants were brought to pH 8 with NaOH. EDTA was added to a final concentration of 10 mM and the supernatant pH was readjusted to 8. Supernatants were filtered (0.45 µm, Millipore) and loaded on a protein A column (protein A sepharose 4 fast flow or HiTrap protein A, both from Pharmacia). The column was washed with PBS containing 0.5 M NaCl until the absorbance at 280 nm decreased to baseline. The column was washed in PBS and eluted with 0.5 M acetic acid. Column fractions were immediately neutralized by eluting into tubes containing 1 M Tris pH 9. The peak fractions containing the MuSK RB were pooled and dialyzed versus PBS. Recombinant Autographa californica baculovirus encoding the Dmk (MuSK) RB was designated "vDmk receptorbody" and deposited with the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland 20852 on May 16, 1995 under ATCC Accession No. VR-2507.

EXAMPLE 5 - SEQUENCING OF HUMAN MuSK RECEPTOR

5 In order to obtain the full coding sequence of the human MuSK receptor, oligonucleotides based on the rat sequence were utilized as PCR primers to amplify cDNA from a human muscle biopsy. The PCR fragment so produced was then sequenced and the resulting new sequence corresponded to a partial sequence of the human MuSK receptor. The novel partial human MuSK receptor sequence was then used to obtain further sequence through
10 successive rounds of the RACE procedure. (Frohman, M. A. (1990), RACE: Rapid amplification of cDNA ends. in PCR Protocols, Innis, M.A.Gelfand, D.H., Sninsky, J.J., and White, T.J. eds. Academic Press. San Diego).

15 This process was complemented by obtaining human genomic clones of MuSK and using the coding sequence of the genomic MuSK to design oligonucleotide primers used to amplify the biopsy cDNA. Stretches of the human MuSK cDNA sequence which were difficult to sequence, absent or presenting some ambiguity were confirmed, corrected or added from the human genomic MuSK sequence. MuSK cDNA variants produced by
20 alternative splicing of MuSK transcripts may be obtained by using this sequence to obtain MuSK cDNA from human sources. The deduced amino acid sequence of the human MuSK receptor and the nucleotide sequence encoding it is set forth in Figure 4. One of skill in the art will readily recognize that this sequence may be used to clone full length, naturally
25 occurring cDNA sequences encoding the human MuSK receptor, which may vary slightly from the sequence set forth in Figure 4.

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EXAMPLE 6 - HOMOLOGOUS RECOMBINATION TO DISRUPT THE
MuSK GENE

5 The tyrosine kinase domain of MuSK is comprised of 11 subdomains that are divided among three coding exons. Subdomain I encoding the ATP-binding domain is located on the first kinase exon, while subdomains 5-11 encoding the catalytic region are located on the third kinase exon (Figure 5). To disrupt MuSK tyrosine kinase activity, a targeting vector was designed that would
10 delete most of the third kinase exon upon homologous recombination into the endogenous mouse MuSK locus (Figure 5); this targeting vector contained a total of 3.8 kb of homology with the mouse MuSK gene.

The MuSK gene targeting vector was constructed from mouse genomic DNA
15 fragments isolated from a lambda FIX II phage library prepared with 129 strain mouse genomic DNA (Stratagene). The 1.7 kb SpeI fragment depicted in Figure 5 was ligated into the compatible ends of a unique XbaI site upstream of the PGK-neo cassette (destroying the SpeI and XbaI sites), while the 2.1 kb BamHI DNA fragment depicted in Figure 5 was blunt-end ligated into the
20 unique HindIII site between the PGK-neo cassette and MC1-tk expression cassettes (destroying the BamHI and HindIII sites). The targeting vector was linearized by digestion with NotI and then electroporated into E14.1 embryonic stem cells, which were subjected to a double selection protocol (gancyclovir addition resulted in a 5-10 fold enrichment compared with
25 selection in G418 alone) and then used to generate chimeric mice as previously described (Conover et al., 1995; DeChiara et al., 1995).

Successful gene targeting using this construct was predicted to result in the generation of a novel 3.8 kb EcoRI fragment from the targeted allele as
30 detected by a 5' probe, as well as loss of two NcoI fragments hybridizing to a kinase probe (Figure 5). Southern blot screening for these fragments revealed

that successful targeting of the mouse MuSK gene was achieved in four of approximately 400 embryonic stem (ES) cell clones obtained using a double selection scheme intended to enhance for selection of targeted clones; the ES cells were derived from the 129 strain of mice. Male chimeras derived from all four of these targeted clones were bred with C57BL/6 females. Chimeras from two of the targeted clones transmitted the mutant allele to the F1 generation. The resulting F1 progeny heterozygous for the MuSK mutation were viable and appeared normal and fertile.

EXAMPLE 7 - MuSK GENE DISRUPTION RESULTS IN PERINATAL
LETHALITY

The heterozygous F1 progeny were interbred to generate mice homozygous for the MuSK gene disruption (designated MuSK^{-/-} mice). Among the F2 litters derived from these crosses were newborn mice that died perinatally. Genotype analysis of tail DNA mice revealed that the dead pups were homozygous for the mutant MuSK allele (Figure 6); significantly, not a single mouse homozygous for the mutation survived the perinatal period (37 homozygotes were noted among the first 138 pups that were genotyped, corresponding to a 26.8% frequency of homozygotes).

To determine the phenotype of the MuSK^{-/-} newborns immediately at birth, applicants were careful to observe the births of several litters derived from heterozygote crosses. Though normal in their gross anatomy and body weight, the MuSK^{-/-} pups differed in several striking ways from their littermate controls. First, they showed no spontaneous movement and did not respond to a mild tail or leg pinch. Only a strong tail pinch was able to elicit a weak uncoordinated movement. By contrast, littermate controls showed extensive movement and responded vigorously to a mild tail pinch. Second, the MuSK^{-/-} pups were cyanotic at birth and appeared not to breathe, although their hearts continued to beat for a short time after birth. To determine

whether the MuSK^{-/-} pups had ever taken a breath, applicants examined the lungs histologically. Lung alveoli are collapsed in utero, and expand with the first breath of life; even if respiration is then terminated, the alveoli remain expanded.

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Histological examination (Figure 7A) revealed that the alveoli of MuSK^{-/-} pups were not expanded, indicating that the pups had never taken a breath. In contrast, the lungs of the littermate controls displayed expanded alveoli (Figure 7B).

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EXAMPLE 8 - NORMAL SKELETAL MUSCLE IN MuSK^{-/-} MICE

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Because MuSK is localized to synaptic sites in skeletal muscle (Valenzuela, D., et al., 1995, Neuron 15: 573-584) and because MuSK^{-/-} mutant mice are immobile at birth and die shortly thereafter, applicants reasoned that neuromuscular synapse formation might be aberrant in MuSK^{-/-} mutants. Applicants first examined the diaphragm muscle because its simple organization and thin structure allows synaptic sites to be visualized in whole-mount preparations. The diaphragm muscle is innervated by the phrenic nerve, which normally enters near the center of the diaphragm muscle. The main intramuscular nerve is oriented perpendicular to the long axis of the muscle fibers and extends through the central region of the muscle.

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For whole-mount diaphragm preparations newborn mice were fixed in 1% paraformaldehyde in phosphate-buffered saline (PBS) at 4°C for several hours and then rinsed briefly in PBS. Diaphragm muscles were dissected out, washed twice for 10 minutes in PBS, incubated in 0.1M glycine in PBS for 15 minutes, rinsed for 5 minutes in PBS, and permeabilized with 0.5% Triton X-100 in PBS (PBT) for 5 minutes. The muscles were then incubated with rabbit antibodies to synaptophysin (kindly provided by Dr. R. Jahn, Yale University Medical School), which were diluted 1/1000 in PBT with 2% BSA,

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overnight at 4°C, subsequently rinsed for 5 minutes in PBT, washed three times for one hour in PBT and then incubated simultaneously with fluorescein-conjugated sheep anti-rabbit IgG (Boehringer Mannheim) and tetramethylrhodamine-conjugated α -bungarotoxin (α -BGT) (Molecular Probes, Oregon) overnight at 4°C. The tissues were then washed three times for 1 hour with PBT, rinsed once with PBS for 5 minutes, fixed in 100% methanol at -20°C and mounted in 90% glycerol, 0.1 M Tris, pH 7.5 with 1 mg/ml *p*-phenylenediamine. The whole-mounts were viewed with epifluorescence and filters that were selective for rhodamine or fluorescein, and images were recorded either on film or on a CCD camera (Princeton Instruments).

The arrangement and gross structure of the muscle fibers (compare Figure 7C and 7D), as well as of the main intramuscular nerve, appeared to be unaltered in MuSK^{-/-} mutant mice. Thus, although the onset of MuSK expression occurs at about embryonic day 11 in developing mouse somites (within the presumptive myotome), MuSK does not appear to be essential for the generation, proliferation and fusion of myoblasts, or for the growth of motor axons from spinal cord to muscle.

EXAMPLE 9 - AGRIN FAILS TO INDUCE AChR CLUSTERING IN MYOTUBES LACKING MuSK

The localization of MuSK to the NMJ inspired us to ask whether MuSK is required for responsiveness to agrin. To test this, applicants first isolated myoblasts from newborn MuSK^{-/-} mice or from control pups, attempted to differentiate them into myotubes in culture, and then assayed for their responsiveness to agrin.

Primary myoblast cultures were established from hind limb musculature of newborn MuSK^{-/-} or littermate control pups. This tissue was treated sequentially with collagenase and trypsin, then plated onto plastic tissue culture dishes. After 1 hour, non-adherent cells (principally myoblasts) were removed and plated onto chamber slides coated with poly-D-lysine and fibronectin. Myoblast cultures were maintained in Dulbecco's Modified Eagle's Medium (DMEM) containing 25% fetal calf serum, 10% horse serum, and 50 µg/ml gentamycin. To induce myotube formation, cultures were switched to a medium consisting of DMEM containing 5% horse serum, L-glutamine and gentamycin to which 20 µM cytosine arabinoside was added after 24 hr. After an additional 2-3 days, contractile myotubes had formed abundantly in cultures from both MuSK^{-/-} and control pups. C2C12 cells were maintained and caused to differentiate in a serum-poor medium as previously described (Ferns, M., et al., 1993, Neuron 11: 491-502).

For agrin-mediated AChR clustering assays on primary myotubes, cultures on chamber slides were treated overnight with c-agrin_{4,8} at 0.01-100 nM; for evaluating MuSK-Fc as an inhibitor of clustering, differentiated C2C12 cells, on chamber slides coated with fibronectin and poly-D-lysine, were pretreated with MuSK-Fc or a control receptor-body for 1 hr at 37°C before addition of approximately 10 nM agrin_{4,8} for overnight incubation. Following overnight treatments with agrin, the cells were next incubated in rhodamine-conjugated α-bungarotoxin to label AChRs, then fixed and mounted for fluorescence microscopy. To quantify the extent of AChR clustering, randomly selected myotubes were viewed under fluorescein optics, then switched to rhodamine optics and the number of AChR clusters within a reticule grid aligned along the long axis of the myotube were counted. AChRs on the surface of cultured primary myotubes were quantitated by incubating live cultures with 25mCi 125I-α-BGT for 1 hr at room temperature, washing, and then lysing the cells in 0.1 N NaOH. The protein concentration in aliquots of each extract was

determined using a BCA protein assay kit (Pierce), while the remainder of the extract was counted in a gamma counter.

5 Myoblasts from both the control and MuSK^{-/-} mice were able to fuse and form long, twitching myotubes in culture. Together with the observation that skeletal muscle appears rather normal in MuSK^{-/-} mice, these findings indicate that MuSK is not critical for early muscle development and myoblast fusion. On the other hand, MuSK appeared to be absolutely required for AChR clustering in response to agrin. After stimulation with the most active form of c-agrin, containing both the four and eight amino acid insertions (c-agrin_{4,8}), AChR clusters were evident only in the myotubes from control mice (Figure 8A). While clusters were induced in normal myotubes with as little as 1 nM c-agrin_{4,8}, no clustering was observed in MuSK^{-/-} myotubes even after increasing the concentration of c-agrin_{4,8} to as high as 100nM (Figure 8B). Lack of detectable clustering was not due to the absence of AChRs, since myotubes from MuSK^{-/-} mice expressed similar numbers of AChR on their surface as did myotubes from control mice (Figure 8C). Thus MuSK appeared to be absolutely required for AChR clustering in response to agrin.

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EXAMPLE 10 - AGRIN INDUCES PROMINENT AND RAPID TYROSINE PHOSPHORYLATION OF MuSK

25 The inability of agrin to induce AChR clustering in myotubes from MuSK^{-/-} mice demonstrates that MuSK is required for agrin responsiveness, and is consistent with the possibility that MuSK serves as the functional agrin receptor. However, since clustering occurs over a period of hours, these results are also consistent with the possibility that MuSK acts much further downstream in the agrin signaling pathway. To begin to distinguish between

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these possibilities, applicants took advantage of the fact that RTKs become rapidly autophosphorylated on tyrosine upon challenge with their cognate ligand. Applicants decided to assay four of the known forms of soluble agrin - which exhibit differing AChR clustering activities (Ruegg, M.A. et al., 1992, 5 Neuron 8: 691-699; Ferns, M., et al., 1992, Neuron 8: 1079-1086; Ferns, M., et al., 1993, Neuron 11: 491-502; Hoch, W. et al., 1994, EMBO J. 13: 2814-2821) - for their ability to induce phosphorylation of the MuSK receptor.

The ability of various agrins and growth factors to induce MuSK or ErbB3 10 tyrosine phosphorylation, for the indicated times and at the indicated concentrations, was evaluated in primary rat myoblasts and in either untransfected C2C12 myoblasts, or in C2C12 myoblasts stably transfected with a chick MuSK-expressing plasmid. The cells were challenged at confluence in an undifferentiated state, or approximately 4-5 days after being induced to 15 differentiate into myotubes in serum-poor media. After challenge, the cells were lysed, the extracts subjected to immunoprecipitation with receptor-specific antibodies, and then immunoblotted with either receptor-specific or phosphotyrosine-specific antibodies, using methods previously described (Stitt, T., et al., 1995, Cell 80: 661-670). Polyclonal 20 antibodies for MuSK were generated as follows: for rat MuSK, rabbits were immunized with a peptide corresponding to the carboxy-terminal 20 amino acids of the rat MuSK protein (Valenzuela, D., et al., 1995, Neuron 15: 573-584; the nomenclature for this antibody is: 41101K); for chick MuSK, rabbits were immunized with a peptide corresponding to the first 19 amino acids of the 25 chick MuSK cytoplasmic domain (Peptide: TLPSELLLDRLHPNPMYQ; the nomenclature for this antibody is 52307K). The specificity of the antibodies was determined on Cos-cell expressed MuSK proteins, by both immune-precipitation and Western, comparing untransfected Cos cell lysates to lysates from rat and chicken-MuSK transfected Cos cells. 41101K immune 30 precipitates and Westerns rodent MuSK, but does not recognize chicken

MuSK. 52307 immune precipitates and Westerns chicken MuSK. Antibodies to ErbB3 were obtained from Santa Cruz Biotechnology, Inc.

5 Cultures of confluent C2C12 cells, either undifferentiated or differentiated in serum-poor media for four to five days as described above, were transferred to 4°C and incubated for 90 minutes with either MuSK-Fc or TrkB-Fc (at 5 mg/ml), each in the presence of the indicated mock or agrin-containing conditioned media (with 100nM agrin). Agrin levels were determined by Western analysis of the conditioned media with a rat agrin antibody (131,
10 from StressGen, Inc.), using a purified agrin control of known concentration. Following these incubations, the cells were washed four times with PBS containing calcium and magnesium, and then incubated for an additional hour with radio-iodinated goat anti-human IgG (NEN/Dupont; 1 mCi/ml in PBS) to detect surface-bound receptor-Fc. After four additional washes, cells
15 were solubilized in 0.1N NaOH, and bound radioactivity was determined. The assay is similar to that described elsewhere (Davis, S., et al., 1994, Science 266: 816-819).

Transient transfections using either previously described agrin constructs
20 (Ferns, M., et al., 1993, Neuron 11: 491-502) or empty vector controls, or stable transfections of a chick MuSK-expression construct, were performed as described (Glass, D., et al., 1991, Cell 66: 405-413; Ip, N.Y., et al., 1992, PNAS (USA) 89: 3060-3064). Agrin concentrations in conditioned media derived from transient transfections were estimated by immunoblot comparisons
25 with purified agrin of known concentration.

Phosphorylation was assessed on the endogenous MuSK receptor that is highly expressed in myotube cultures, obtained by differentiating either the C2C12 mouse myoblast cell line (Valenzuela, D., et al., 1995, Neuron 15: 573-
30 584) or primary rat myoblasts. Strikingly, soluble agrins containing the eight amino acid insert at position Z (c-agrin_{4,8} and c-agrin_{0,8}), which are the forms

capable of inducing AChR clustering, were also the forms that induced prominent tyrosine phosphorylation of MuSK (Figure 9A). The agrin most active in clustering (c-agrin_{4,8}) was also most active in inducing MuSK phosphorylation (Figure 9A). In contrast, the soluble agrins lacking the eight amino acid insert (c-agrin_{4,0} and c-agrin_{0,0}), which cannot induce AChR clustering, also could not induce MuSK phosphorylation (Figure 9A).

The specificity of action of agrin was further explored by comparing its activity to growth factors known to have receptors on muscle. Of the several such factors tested, including insulin, fibroblast growth factor (FGF) and ARIA/neuregulin, only agrin could induce phosphorylation of MuSK (Figure 9A); since FGF also induces AChR clustering on myotubes (Peng, H.B., et al., 1991, Neuron 6: 237-246), these results also indicate that MuSK phosphorylation is specific to agrin responses and not just to agents capable of inducing clustering. Furthermore, while such factors could be shown to induce phosphorylation of their own RTKs on myotubes (e.g., neuregulin induces phosphorylation of its cognate RTK, erbB3), agrin could only activate MuSK and not other RTKs (Figure 9B).

The activation of a RTK by its cognate ligand typically tends to occur rapidly, and applicants could demonstrate that agrin induces tyrosine phosphorylation of MuSK with kinetics similar to those seen for well-characterized RTK/ligand systems (e.g. Kaplan, D.R., et al., 1991, Nature 350: 158-160); induction was detectable by one minute, peaked within the first five minutes, and remained elevated for over an hour (Figure 9D). The tyrosine phosphorylation of MuSK also occurred using agrin at concentrations similar to those noted for other ligands that act on RTKs (Ip, N.Y., et al., 1993, Neuron 10: 137-149), with phosphorylation detectable using 1 nM agrin (Figure 9C).

The requirement of MuSK for agrin responsiveness, the ability of agrin to induce rapid and prominent MuSK phosphorylation, the specificity of agrin for MuSK as compared to other factors tested, and the precise correlation of agrin forms active in AChR clustering assays and in MuSK phosphorylation assays, together continue to support the notion that MuSK serves as the functional agrin receptor.

EXAMPLE 11 - AGRIN DOES NOT DIRECTLY BIND TO AN ISOLATED
MuSK ECTODOMAIN

10 If MuSK is indeed the functional agrin receptor, applicants would expect to be able to demonstrate binding of agrin to MuSK. In an attempt to demonstrate such binding, applicants first constructed an expression construct encoding a fusion protein between the ectodomain of rat MuSK and the Fc portion of human immunoglobulin G1 (designated MuSK-Fc), and then produced and purified the fusion protein. Similar receptor-Fc fusions have previously been used to characterize binding between RTKs and their ligands (Davis, S., et al., 1994, Science 266: 816-819; Stitt, T., et al., 1995, Cell 80: 661-670).

20 Baculovirus expression vectors encoding MuSK-Fc, TrkB-Fc, and Ret-Fc produced fusion proteins in which the ectodomains of rat TrkB, rat Ret, or rat MuSK, respectively, were linked to a spacer with the sequence Gly-Pro-Gly, followed by the hinge, CH2, and CH3 regions of human IgG1, beginning with the residues Glu-Pro-Lys, as described (Davis, S., et al., 1994, Science 266: 816-819). Baculovirus infections into *Spodoptera frugiperda* SF-21AE insect cells were performed by standard methods (Stitt, T., et al., 1995, Cell 80: 661-670). The soluble Fc-containing proteins were purified by protein A-Sepharose (Pharmacia) chromatography.

30 The binding of agrin to immobilized MuSK-Fc as compared to a monoclonal antibody specific for agrin was evaluated by use of BIAcore biosensor

technology (Pharmacia Biosensor), using approaches previously described (Stitt, T., et al., 1995, Cell 80: 661-670). Heparin and CaCl₂ were supplied by Sigma Chemical Co. (St. Louis, MO) and used without further purification. The agrin-specific monoclonal antibody (clone AGR131 generated to rat agrin) was purchased from StressGen Biotechnologies Corp. (Victoria, BC, Canada).

In a first approach, applicants used MuSK-Fc together with BIAcore biosensor technology. The BIAcore technology allows for the direct and quantitative measure of binding of soluble ligands to receptors coupled onto a sensor chip. Recombinant MuSK-Fc was covalently coupled to a surface on the BIAcore sensor chip, and as a control, a monoclonal antibody specific for rat agrin was also coupled to a separate surface on the sensor chip; media containing c-agrin_{4,8} was then passed over the two surfaces. While robust binding of the agrin to the antibody surface was easily detected, no binding of the agrin to the MuSK surface could be seen (Figure 10A). Furthermore, while binding to the antibody surface was specifically competable by excess soluble antibody added to the agrin-containing media, the binding was not competable by excess soluble MuSK-Fc (Figure 10A). Since agrin activity requires calcium (Bowe and Fallon, 1995, Ann. Rev. Neurosci. 18: 443-462), and because some heparin-binding factors require heparin to bind to their receptors (Goldfarb, M., 1990, Cell Growth & Differentiation 1: 439-445), applicants also attempted binding in the presence of calcium or heparin; in neither case was binding to the MuSK surface observed (Figure 10A).

Next, applicants tried to demonstrate binding of MuSK and agrin by attempting to use MuSK-Fc to detect agrin immobilized onto nitrocellulose. In contrast to our control experiments, in which immobilized brain-derived neurotrophic factor (BDNF) was easily detected by an Fc fusion of its cognate receptor (TrkB-Fc), and in which immobilized agrin was easily detected by the agrin-specific monoclonal antibody, immobilized agrin could not be detected by MuSK-Fc (Figure 10B).

The negative binding results described above demonstrate that the isolated MuSK receptor is not sufficient to bind agrin. Thus, despite the plethora of functional data indicating that agrin acts via MuSK, MuSK may not directly serve as a receptor for agrin. Alternatively, MuSK may require additional components or modifications which are required for it to bind and respond to agrin.

10 EXAMPLE 12 - AGRIN ACTIVATES MuSK IN A CELL-CONTEXT-
DEPENDENT FASHION: REQUIREMENT FOR A
MYOTUBE-SPECIFIC ACCESSORY COMPONENT

Based on the results described above, applicants considered the possibility that the agrin-MuSK interaction requires additional components. To further explore this possibility, applicants determined the cell-context dependency for agrin activation of MuSK, reasoning that if an accessory component was required, it might be specifically expressed only on cells normally responding to agrin. Thus applicants ectopically expressed full-length cDNAs encoding rat, human and chicken MuSK in fibroblasts, and assayed for whether these MuSK receptors could be inducibly phosphorylated by agrin. When expressed in fibroblasts, none of the three species of MuSK could be phosphorylated in response to agrin. While this supported the possibility that MuSK requires an accessory myotube-specific component to respond to agrin, it was also possible that our cDNAs encoded MuSK variants that could not respond to agrin. This was a potentially worrisome possibility since there are multiple differently spliced versions of the MuSK transcript (Valenzuela, D., et al., 1995, Neuron 15: 573-584), applicants did not know which of the forms were normally agrin-responsive, and our cDNAs only accounted for a subset of the variant forms. Thus applicants decided to express our cDNAs in myoblasts to verify that they could mediate responses to agrin when expressed in the right

context. For this purpose applicants chose to express the chicken MuSK in the mouse C2C12 myoblast cell line, since the chicken MuSK could easily be distinguished from the endogenous mouse MuSK based on size and by using particular antibodies. When expressed in undifferentiated myoblasts, the chicken MuSK did not undergo phosphorylation in response to any isoforms of agrin (Figure 11, see lanes indicated "Undif", upper panel), just as it did not undergo phosphorylation in fibroblasts; undifferentiated C2C12 cells do not express appreciable amounts of endogenous MuSK (Figure 11, lanes indicated "Undif", lower panel and also (Valenzuela, D., et al., 1995, Neuron 15: 573-584), so applicants could not compare activation of the endogenous mouse MuSK in myoblasts. Upon differentiation into myotubes, the introduced chicken MuSK was as effectively activated by agrin as was the endogenous mouse MuSK (Figure 11, lanes indicated "Diff", upper panel); both introduced and endogenous MuSK had identical profiles of responsivity to the various forms of agrins, with activations mediated only by forms having the eight amino acid insert at the Z position. Thus our cDNAs encode MuSK proteins that are perfectly competent to undergo agrin-induced phosphorylation, but they can only be activated by agrin in the context of a differentiated myotube, consistent with the notion that agrin activation of MuSK requires a myotube-specific accessory component that is not expressed in fibroblasts or undifferentiated myoblasts.

EXAMPLE 13 - A RECEPTOR COMPLEX CAN BE DEMONSTRATED
BETWEEN AGRIN, MuSK AND A MYOTUBE-SPECIFIC
ACCESSORY COMPONENT

Altogether, the above data indicate that agrin requires MuSK to mediate clustering and that agrin activates MuSK very rapidly, but that agrin does not directly bind to a purified MuSK ectodomain and can only activate MuSK in the context of a myotube. These findings are consistent with the possibility

that MuSK is a requisite part of an agrin receptor complex, but that although MuSK provides a key signaling function for this complex, it requires another component(s) to bind to agrin. Similar types of receptor complexes have been described for other ligands. Perhaps some of the best characterized examples

5 include the receptor complexes for ciliary neurotrophic factor (CNTF) and its cytokine relatives (Davis, S., et al., 1993, Science 260: 1805-1808; Stahl and Yancopoulos, 1993, Cell 74: 587-590). In order to interact with its two signal transducing "b" receptor components, gp130 and LIFRb, CNTF must first bind to its "a" receptor component, known as CNTFRa. CNTFRa serves no

10 signaling role, and is in fact linked to the surface via a glycosylphosphatidylinositol linkage and thus has no cytoplasmic domain. The receptor complex for CNTF is built in step-wise fashion: CNTF first binds to CNTFRa; this initial complex can then bind to and recruit a single "b" component; finally, a complete complex forms that involves "b" component

15 dimerization, which is required for signal initiation (Figure 12A). In the final complex, CNTF seems to make contacts with all three receptor components. Interestingly, receptor complexes for CNTF can be built in solution using just the soluble ectodomains of the various components. Furthermore, if just one of the receptor components is linked to the surface, a receptor complex can be

20 built around it using soluble versions of the other components, but only in a CNTF-dependent fashion (Figure 12B).

If agrin binds to MuSK in a receptor complex, applicants reasoned that they might be able to manipulate this complex in much the same way the CNTF

25 receptor complex can be manipulated. To explore the possibility that myotubes specifically express an accessory component(s) required for agrin to bind MuSK (Figure 12C), applicants decided to test whether applicants could specifically build a receptor complex on the surface of myotubes, but not on other cells, using agrin together with a soluble version of the MuSK receptor

30 to complex to the putative accessory component(s) on the surface of myotubes. Confirming this possibility, applicants found that the binding of

soluble MuSK-Fc to the surface of cells can be increased using agrin, but only on the surface of differentiated myotubes and not on the surface of fibroblasts or myoblasts (Figure 13A). These data demonstrate that complexes can form between agrin and MuSK, but only in the presence of a myotube-specific component(s) (as suggested in Figure 12C). Interestingly, although forms of c-agrin containing the eight amino acid insert at the Z position are best able to promote agrin-dependent MuSK complex formation, forms of c-agrin without this insert can also form these complexes. The ability of all the soluble forms to promote complex formation, including those lacking the eight amino acid insert for activity, may be related to previous findings that matrix-bound forms of agrins lacking the Z insert can activate clustering (Ferns, M., et al., 1992, Neuron 8: 1079-1086). Thus although soluble agrins lacking inserts at the Z position do not seem capable of signaling, they may be able to form partial complexes, while matrix-associated forms of these same agrins can proceed to form complete signaling-competent complexes. Interestingly, ligands for the EPH family of RTKs provide an example of ligands that bind but do not activate their receptors when presented in soluble form, but which can act as potent activators when bound to the cell surface (Davis, S., et al., 1994, Science 266: 816-819); deliberate clustering of the soluble ligands can allow them to activate as well, suggesting that the role of surface-attachment is to allow for ligand-clustering (Davis, S., et al., 1994, Science 266: 816-819).

In the absence of added agrin, the MuSK-Fc exhibited much higher binding to myotube surfaces than did several control receptor-Fc fusion proteins (Figure 13A, data shown for TrkB-Fc); the MuSK-Fc, however, displayed similar agrin-independent binding to both myoblast and fibroblasts as did control receptor-Fc proteins (Figure 13A). Specific binding of MuSK-Fc to myotube surfaces, in the absence of exogenously provided agrin, may indicate that MuSK has an affinity for its myotube-specific accessory component in the absence of ligand. Alternatively, since myotubes make muscle forms of agrin

(lacking the eight amino acid insert at the Z position), the specific binding of MuSK-Fc to myotubes in the absence of added agrin could be explained by the formation of a complex between the added MuSK-Fc and endogenously expressed muscle agrin along with the accessory component; adding
5 additional exogenous soluble agrin may simply allow for even more MuSK to be recruited into complexes with the myotube-specific accessory component. Although both myoblasts and myotubes make endogenous agrin, myoblasts seemingly cannot form complexes with added MuSK-Fc since they do not express the required accessory component.

10

To confirm that MuSK directly interacts with agrin as part of its receptor complex, applicants next demonstrated that radiolabelled agrin could be cross-linked to MuSK receptors on the surface of myotubes.

15

Flg-tagged human agrin protein corresponding to the COOH-terminal 50 kD of human agrin 4,8 was expressed in Cos cells and purified by affinity and size-exclusion chromatography to >95% purity. Twenty µg were iodinated by a modification of the lactoperoxidase method described previously (DiStefano, P., et al, 1992, Neuron 8: 983-993). Incorporation of ¹²⁵I was greater than 80%;
20 ¹²⁵I-h-agrin 4,8-flg was separated from free ¹²⁵I on a 1x3 cm Sephadex G-25 column prior to use in cross-linking assays. Specific activity was ~4000 cpm/fmol (~2400 Ci/mmol). Biological activity of ¹²⁵I-h-agrin 4,8-flg was monitored by tyrosine phosphorylation of MuSK in C2C12 myotubes and was found to be indistinguishable from its unlabeled counterpart. For
25 cross-linking studies, 10 cm plates of differentiated C2C12 myotubes were incubated in 1 nM of [¹²⁵I]-agrin_{4,8} in 1.5 ml of PBS containing 1% BSA and 1 mg/ml glucose in the presence or absence of 150-fold excess unlabeled agrin_{4,8} for 75 min at 4°C. The cross-linking agent DSS (disuccinimidyl suberate) was added to a final concentration of 0.2mM and the plates were incubated at
30 room temperature for 30 min, washed 3 times with 50 mM Tris/150mM NaCl pH 7.5, lysed, and subjected to immunoprecipitation with MuSK-specific

antibodies. For peptide competition, peptide antigen was included in the immunoprecipitation at a final concentration of 20 µg/ml. The samples were then electrophoresed and the fixed and dried gels were exposed for autoradiography.

5

Immunoprecipitations using a MuSK-specific antibody, from lysates of myotubes chemically cross-linked to radiolabelled recombinant human agrin contained complexes corresponding in size to agrin/MuSK complexes (Figure 13B). These agrin/MuSK complexes were not seen in the presence of excess
10 unlabelled agrin, or if a peptide was used to block MuSK precipitation (Figure 13B). Additional radiolabelled species that immunoprecipitated with the MuSK antibody correspond to forms of agrin that are associated with, but not cross-linked to, MuSK, presumably due to the low efficiency of cross-linking (Figure 13B); low levels of additional agrin complexes, perhaps involving
15 MASC, could also be detected in these immunoprecipitations.

Finally, if our findings that soluble MuSK could form complexes with its requisite myotube-specific accessory component are correct, then this soluble receptor should also act as an inhibitor of agrin-mediated responses by
20 sequestering the accessory component and preventing it from interacting with the endogenously-expressed, signaling-competent MuSK. Indeed, addition of increasing amounts of MuSK-Fc did inhibit agrin-mediated clustering of AChRs (Figure 13C) as well as agrin-induced MuSK phosphorylation in a dose-dependent manner, while control receptor-Fc proteins had no inhibitory
25 effect.

EXAMPLE 14 - CLONING OF HUMAN AGRIN cDNA

30 Probes corresponding to human agrin were prepared by PCR based on partial sequences of human agrin available from the Genbank database.

Two pairs of PCR primers were synthesized based on human agrin cDNA sequences obtained from Genbank. The sequences of the oligonucleotide primers were as follows:

5 Primer pair 18:

h-agrin 18-5' : 5'-GACGACCTCTTCCGGAATTC-3'

h-agrin 18-3' : 5'-GTGCACATCCACAATGGC-3'

Primer pair 35:

10 h-agrin 35-5' : 5'-GAGCAGAGGGAAGGTTCCCTG-3'

h-agrin 35-3' : 5'-TCATTGTCCCAGCTGCGTGG-3'

The oligonucleotide primers were used for PCR amplification of two segments of DNA of approximately 100 nts (primer pair 18) and 85 nts (primer pair 35) using 300 ngs of human genomic DNA as a template. The PCR amplification was carried out as recommended by the manufacturer (Perkin-Elmer) under the following conditions: 35 cycles at 94°C for 60 sec, 55°C for 50 sec and 72°C for 30 sec. The PCR fragments obtained were purified from an agarose gel and re-amplified for 30 cycles using the same PCR conditions described above.

After amplification, the PCR reactions were electrophoresed in agarose gels, the agarose containing the DNA bands of 100 and 85 nts respectively was excised, purified by QiaEx II (Qiagen), and then cloned into plasmid pCR-script using Stratagene's pCR-Script cloning kit, followed by bacterial transformation and plating onto agar-ampicillin plates as recommended by the manufacturer. Bacterial colonies containing the 100 and 85 nt inserts were identified by PCR using the primers described above. The PCR fragments obtained were radiolabeled for use as probes using a standard PCR reaction (Perkin-Elmer) on 20 ng of DNA template, except that 5 nmoles each of dATP, dGTP and dTTP and 0.2 mCurie of alpha ³²P-dCTP (Du Pont 3000 Ci/mmol)

were added to the reaction mixture and then subjected to 7 cycles of PCR.

Unincorporated label was separated from the probes on a G50 NICK column (Pharmacia). These probes were used to screen a human fetal brain cDNA library (Stratagene Cat# 936206) using standard library screening procedures (Sambrook, Fritsch and Maniatis, Molecular Cloning, a Laboratory Manual, (1989) Second Edition, Cold Spring Harbor Laboratory Press). One and a half million phage plaques were plated in XL-1 Blue bacteria as recommended by Stratagene, and transferred to nitrocellulose filters in duplicate as previously described (*Id.*). The filters were processed and each filter replica was used for hybridization with one of the above probes as previously described (*Id.*).

Plaques hybridizing to both probes were isolated and purified and a plasmid containing the cDNA insert was excised from the lambda clone according to Stratagene's recommended procedure (EXASSIST/SOLR System). The pBluescript plasmid containing the human Agrin insert was purified and the insert was then sequenced using an automated sequencing kit (Applied Biosystems).

As a result of this screen, one clone (pBL-hAgrin1) was obtained which contains a nucleotide sequence encoding an amino acid sequence of human agrin. The first amino acid encoded by the cloned nucleotide sequence corresponds approximately to amino acid 424 of rat agrin (See Figure 14). The nucleotide sequence of the clone ends downstream of the stop codon. Clone pBL-hAgrin1 contains a 4 amino acid insert starting at the position which corresponds to position 1643 of Figure 14, a point which was previously described for the rat as position "Y" (Stone, D.M. and Nikolics, K., J. Neurosci. 15: 6767-6778 (1995)). The sequence of the 4 amino acid insert both in clone pBL-hAgrin1 and in the rat is KSRK.

A second clone was obtained from this screen. This second clone (pBL-hAgrin23) also contains a nucleotide sequence encoding an amino acid sequence of human agrin. The first amino acid encoded by the cloned

nucleotide sequence corresponds approximately to amino acid 1552 of the rat agrin (See Figure 14). The nucleotide sequence of the clone ends downstream of the stop codon. Clone pBL-hAgrin23 contains an 8 amino acid insert starting at a position which corresponds to position 1780 of Figure 14, a point which was previously described for the rat as position "Z" (Stone, D.M. and Nikolics, K., J. Neurosci. 15: 6767-6778 (1995)). The sequence of the eight amino acid insert both in clone pBL-hAgrin23 and in the rat is ELANEIPV. As previously discussed, it has been reported that the 8 amino acid insert plays an important role in regulating the AChR clustering activity of different agrin forms. Therefore, by inserting a nucleotide sequence encoding the eight amino acid sequence ELANEIPV into clone pBL-hAgrin1 at the position corresponding to position Z of rat agrin, a human 4-8 agrin clone may be obtained. The addition of the 8 amino acid insert at position Z should confer a high level of biological activity to the human 4-8 clone.

Clone pBL-hAgrin23 also contains the 4 amino acid "Y" insert as described above for clone pBL-hAgrin1. However, clone pBL-hAgrin23 contains 17 extra amino acids at the same "Y" position, such that the sequence of the "Y" insert in clone pBL-hAgrin23 is KSRKVLSASHPLTVSGASTPR. Therefore, in addition to the (4-0) and (4-8) human agrin splice variants described above, human clones corresponding to splice variants containing (Y-Z) inserts of (17-0), (17-8), (21-0), and (21-8) are indicated by these results and are within the scope of the present invention.

EXAMPLE 15 - EXPRESSION OF HUMAN AGRIN

Construction of human agrin expression vector

A human agrin Sfi I - Aat II fragment containing the 4 amino acid insert at the position corresponding to the Y-site described for rat agrin (see Figure 14)

was excised from clone pBL h agrin-1. A human agrin Aat II - Not I fragment containing the 8 amino acid insert at the position corresponding to the Z-site described for rat agrin (see Figure 14) was excised from clone pBL h agrin-23. A Xho I - Sfi I fragment was then generated via PCR that contained a
5 preprotrypsin signal peptide, the 8 amino acid flg peptide (from the flag tagging system, IBI/Kodak, Rochester, NY) and the human agrin sequence corresponding to the sequence of amino acids from position 1480 to the Sfi I site located at amino acids 1563-1566 of rat agrin (see Figure 14). The three
10 fragments were then ligated into a Xho I - Not I digested pMT21 expression vector to form the human agrin 4-8 expression vector pMT21-agrin 4-8. The sequence of human agrin 4-8 that was encoded in the expression vector is shown in Figure 15. Expression vectors for the human clones corresponding to splice variants containing (Y-Z) inserts of (0-8) and (4-0) were also constructed.

Expression of human agrin (4-8) in E. coli

The gene for human agrin 4-8 was PCR amplified from pMT21-agrin 4-8 with the primer pair AG5'
20 (5'-GAGAGAGGTTTAAACATGAGCCCCTGCCAGCCCAACCCCTG-3') and AG3' (5'-CTCTGCGGCCGCTTATCATGGGGTGGGGCAGGGCCGCAG-3'). The PCR product was digested with the restriction enzymes Pme I and Not I and cloned into the Pme I and Not I sites of the vector pRG501, a pMB1 replicon that confers kanamycin resistance and is designed to express cloned
25 genes from the phage T7 promoter. One isolate was characterized and named pRG531. The 1315 base pair Nco I - Nae I fragment internal to agrin in pRG531 was then replaced with the corresponding fragment from pMT21-agrin 4-8. The resulting plasmid, pRG451, was transformed into the expression strain RFJ209 [IN(rrnD-rrn/E)1 lacI^Q lacZpL8 fhuAD322-405
30 rpoS_(MC4100) ara::(lacUV5-T7 gene 1)8]. Cultures of RFJ209 / pRG541 induced with IPTG express human agrin to about 5% of total cellular protein and

fractionates with soluble protein upon cell disruption. The crude soluble protein fraction containing human agrin 4-8, as well as human agrin 4-8 purified by Q-Sepharose chromatography was determined to be active in phosphorylation of MuSK receptor.

5

Expression of human agrin (4-8) in *Pichia pastoris*

The 50kD active fragment (portion) of human agrin 4-8 was cloned by PCR using a primer containing a portion of the *S. cerevisiae* α mating factor pre-pro secretion signal and the 5' end of the region encoding the 50kD agrin fragment (GTATCTCTCGAGAAAAGAGAGGCTGAAGCT
10 AGCCCTGCCAGCCCAACC), and a primer containing sequences from the region 3' of the agrin coding region and a NotI site (AATAGTGCGGCCGCGCAACACTCAGGCAAGAAAATCATATC). After PCR
15 the fragment was digested with XhoI, which recognizes sequences in the 5' primer, and NotI, and was cloned into pPIC9 (Invitrogen) digested with XhoI and NotI. The resulting clone was digested with NotI and partially digested with NcoI to remove most of the PCRed agrin sequences. This region was replaced by a NotI-NcoI fragment of agrin from pRG541. PCRed regions were
20 sequenced and shown to be wild-type. This clone, pRG543 was digested with SalI and transformed into *Pichia pastoris* by electroporation. Transformants were selected for a His⁺ Mut⁺ phenotype. Induction of the AOX1 promoter driving the expression of hAgrin was done by growing the cells in buffered glycerol-complex medium containing 0.5% glycerol, pH=6.0, for 24 hrs until
25 the glycerol was exhausted, at which point methanol was added to a final concentration of 0.5%. The culture was centrifuged and the supernatant was dialyzed against PBS. The concentration of hAgrin was approximately 10ug/ml and was determined to be active in phosphorylation of MuSK receptor.

30

Production of Human Agrin 4.8 from Baculovirus Infected Insect Cells

Virus Production

The flg-tagged gene for human agrin 4-8 was engineered into a baculovirus expression plasmid and recombined with viral DNA to generate recombinant baculovirus, amplified and harvested using methods previously described (O'Reilly, D.R., L.K. Miller, and V.A. Luckow, Baculovirus Expression Vectors - A Laboratory Manual 1992, New York: W.H. Freeman). SF21 insect cells (*Spodoptera frugiperda*) obtained from Invitrogen were adapted and expanded at 27°C in Gibco SF900 II serum-free medium. Uninfected cells were grown to a density of 1×10^6 cells/mL. Cell density was determined by counting viable cells using a hemacytometer. The virus stock for FLAG-agrin was added to the bioreactor at a low multiplicity 0.01-0.1 PFU/cell to begin the infection. The infection process was allowed to continue for 3-4 days allowing maximum virus replication without incurring substantial cell lysis. The cell suspension was aseptically aliquoted into sterile centrifuge bottles and the cells removed by centrifugation (1600 RPM, 30 min). The cell-free supernatant was collected in sterile bottles and stored at 4°C in the absence of light until further use.

The virus titer was determined by plaque assay as described by O'Reilly, Miller and Luckow. The method is carried out in 60mm tissue-culture dishes which are seeded with 1.5×10^6 cells. Serial dilutions of the virus stock are added to the attached cells and the mixture incubated with rocking to allow the virus to adsorb to individual cells. An agar overlay is added and plates incubated for 5 days at 27°C. Viable cells were stained with neutral red revealing circular plaques which were counted to give the virus titer expressed in plaque forming unit per milliliter (PFU/mL).

Infection of Cells for Protein Production

Uninfected SF21 cells were grown in a 60L ABEC bioreactor containing 40L of Gibco SF900 II medium with gentamicin sulfate (25 mg/L) and amphotericin B (1 mg/L). Temperature was controlled at 27C and the dissolved oxygen level was maintained at 50% of saturation by controlling the flowrate of oxygen in the inlet gas stream. When a density of 2×10^6 cells/mL was reached, the cells were concentrated within the bioreactor to 20L, using a low shear steam sterilizable pump and a tangential flow filtration device with Millipore Prostack 0.45 micron membranes. After concentration, fresh sterile growth medium was slowly added to the bioreactor while the filtration system continued to remove the spent growth medium by diafiltration. After two volume exchanges an additional 20L of fresh medium was added to the bioreactor to resuspend the cells to the original volume of 40L.

The amount of virus stock required was calculated based on the cell density, virus titer and the desired multiplicity of infection (MOI). Multiplicity ratios between 1 and 10 pfu/cell have been effectively used. The virus stock was added aseptically to the bioreactor and the infection was allowed to proceed for three to four days.

Recovery and Chromatographic Purification

At the conclusion of the infection phase of the bioreactor process the cells were concentrated in the bioreactor using a 30 ft² Millipore Prostack filter (0.45 micron) pore size. The cell-free permeate passing through the filter was collected in a clean process vessel. The protein was diafiltered into a low conductivity buffer (20 mM citrate, pH 5.5) using Millipore Pellicon ultrafiltration membrane cassettes totaling 20 ft² with a nominal 10 kiloDalton cutoff. The protein in the retentate was loaded onto a cation exchange column (Pharmacia SP Sepharose FF) equilibrated with 20 mM citrate buffer, pH 5.5. After loading the protein was washed first with 20 mM citrate, 200 mM sodium chloride, pH 5.5 then with 20 mM Bicine, pH 8.0 to remove contaminating proteins. The protein was eluted with a 0-750 mM

sodium chloride linear gradient over 7.5 column volumes. The eluted agrin was buffer exchanged into 20 mM Tris, pH 8.5 buffer to remove salt for subsequent binding to an anion exchange column.

5 The agrin was then bound to a Pharmacia Q Sepharose FF column equilibrated with 20 mM Tris, pH 8.5. After loading the column was washed with the same buffer to remove contaminating proteins and the protein eluted with a 0-250 mM sodium chloride gradient. The fractions containing agrin were pooled and concentrated and dialyzed into PBS containing calcium
10 and magnesium.

Expression of human agrin (4-8) in COS-7 cells

Lipofectamine reagent (GIBCO-BRL, Inc.) and recommended protocols were used to transfect COS-7 cells with the human agrin cDNA clone pMT21-agrin
15 4-8 containing a nucleotide sequence encoding the eight amino acid sequence ELANEIPV at the position corresponding to position Z of rat agrin. COS media containing secreted ligand was harvested after three days and concentrated 20-fold by diafiltration (DIAFLO ultrafiltration membranes, Amicon, Inc.). The quantity of active human agrin present in the media was
20 determined and expressed as the amount (in resonance units, R.U.) of MuSK receptor specific binding activity measured by a BIAcore binding assay.

EXAMPLE 16 - PREPARATION OF TRUNCATED MOLECULES 25 CONTAINING THE MUSK ACTIVATING PORTION OF HUMAN AGRIN

It has recently been reported that a 21kD fragment of chick agrin is sufficient to induce AChR aggregation (Gesemann, M., et al., 1995, J. Cell. Biol. 128: 625-636). Applicants therefore decided to investigate the properties of various
30 portions of human agrin and to test the ability of each to induce phosphorylation of the MuSK receptor.

As set forth in Figure 15, the amino acid sequence of the 50 kD active portion of human agrin 4,8 is 492 amino acids long. A preprotrypsin signal sequence (Stevenson et al., 1986. Nucleic Acids Res. 21: 8307-8330) precedes a FLAG tag sequence (Hopp et al. 1988. Bio/Technology 6: 1204-1210); together, they
5 constitute the first 23 amino acids. Thus the agrin 4,8 sequence begins with amino acid 24. Truncated molecules were created, each of which contained the signal sequence and FLAG tag (23 amino acids) followed by the agrin 4,8 sequence to which N-terminal deletions had been made to create portions of
10 agrin (designated delta 3 through 9) as follows:

delta 3: agrin sequence starts with amino acid #60: QTAS...
delta 4: agrin sequence starts with amino acid #76: NGFS...
delta 5: agrin sequence starts with amino acid #126: VSLA...
15 delta 6: agrin sequence starts with amino acid #178: GPRV...
delta 7: agrin sequence starts with amino acid #222: GFDG...
delta 8: agrin sequence starts with amino acid #260: ASGH...
delta 9: agrin sequence starts with amino acid #300: AGDV...
All of the sequences continue to the terminal amino acids PCPTP, as with the
20 50kD agrin.

The truncated molecules were made as follows: PCR primers were designed consistent with the nucleotide sequences encoding the first and last ten amino acids of each construct. Included in the 5' primer was sequence data to append
25 the preprotrypsin signal sequence and "FLAG-tag" to the amino terminus of each agrin fragment. Thus, the shortest truncated molecule (delta 9) contains the signal sequence and FLAG tag and the human agrin sequence from amino acid 300 to 492 of human c-agrin 4,8. DNA encoding the "delta" forms of truncated c-agrin 4,8 was then cloned into a eukaryotic expression vector, and
30 transient transfections were performed as previously described (Glass, D., et al., 1991, Cell 66: 405-413; Ip, N.Y., et al., 1992, PNAS (USA) 89: 3060-3064).

Agrin levels in the COS-transfected supernatants were determined by Western analysis of the conditioned media with an agrin-specific antibody (Stressgen 131), using a purified agrin control of known concentration. Each of the truncated molecules was expressed and shown to be capable of inducing tyrosine phosphorylation of MuSK receptor.

Figure 16 shows that the delta 9 truncated molecule can induce phosphorylation of the MuSK receptor, though with less efficiency than the 50kD agrin 4,8 molecule. Thus, each of the truncated molecules created exhibited the biological activity of human agrin 4,8 with respect to the MuSK receptor.

EXAMPLE 17 - PEGylation of Agrin 4,8 and Pharmacokinetics Study

After intravenous administration, the 50kD agrin 4,8 was cleared rapidly from the systemic circulation with a half-life of <10 minutes. (See Figure 17). It was known that the properties of certain proteins can be modulated by attachment of polyethylene glycol (PEG) polymers, which increases the hydrodynamic volume of the protein and thereby slows its clearance by kidney filtration. (See, e.g. Clark, R., et al., 1996, J. Biol. Chem. 271: 21969-21977). Therefore, in an attempt to increase its circulating half-life, agrin 4,8 was modified by covalent attachment of a polyethylene glycol molecule and the effect on the protein's serum half-life was studied.

A solution of 500 mg of human agrin 4,8 at 3.0 mg/mL was added to a reaction vessel and the PEGylation reaction was initiated by addition of monomethoxypoly-ethyleneglycol succinimidyl propionate (PEG) (approximate molecular weight = 20,000 daltons). A ratio of 1.75 moles PEG reagent to agrin was used for the reaction which was carried out at a pH of 7.3

to 8.5 in phosphate buffered saline over a period of 2 hours. The reaction was stopped by addition of 50 mM Tris hydrochloride.

The PEGylated agrin was diluted with buffer at a pH of 8.2 to lower both the conductivity and the concentration of Tris, before loading onto a cation exchange column (Pharmacia SP High Performance). The column was eluted using a gradient from 0 to 600 mM sodium chloride in Tris buffer at a pH of 8.2. Numerous distinct forms of PEGylated agrin eluted along the gradient and unmodified agrin eluted at the high salt end of the gradient.

Monopegylated forms were selectively pooled for subsequent in vivo testing.

Adult Sprague-Dawley rats (male or female) weighing 300-500 g were anesthetized with ketamine/xylazine (50/10 mg/kg) and the right hind limb muscles were denervated by transecting the sciatic nerve at mid thigh level.

After 10-14 days (when MuSK expression was substantially elevated in denervated muscle) rats were re-anesthetized and the right jugular vein was exposed by cut down surgery. Rats were then administered doses of agrin 4,8 or PEG-agrin 4,8 ranging from 1-10 mg/kg into the jugular vein with a 27 gauge needle. After injection, the wound was sutured and tail blood samples were taken at 0 (pre-injection), 5, 10, 15, 30 minutes, and at 1, 2, 4, 6, 8, 16, 24, and 48 hours after the injection. Serum samples were harvested from centrifuged blood and assayed using an ELISA. Serum levels are expressed as µg/ml of serum.

As shown in Figure 17, at a dose of 10 mg/kg, i.v., agrin 4,8 was rapidly cleared from the blood with a half-life of ~10 minutes. In contrast, the half-life of PEG-agrin 4,8 (also at 10 mg/kg, i.v.) was dramatically increased by ~10-fold. These results show that modification of agrin 4,8 with PEG greatly increases its apparent half-life in the blood. Thus, PEGylated agrin may have prolonged activity on MuSK in denervated skeletal muscle and may thus be a more effective treatment for muscular disorders and conditions such as muscle

atrophy. It is expected that the PEGylation of the above-described truncated agrin molecules would similarly increase their apparent half-lives in the blood.

5

DEPOSIT OF MICROORGANISMS

A clone designated pBluescript SK-containing dmk was deposited with the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland 20852 on July 13, 1993 under ATCC Accession No. 75498.

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Recombinant Autographa californica baculovirus encoding the rat Dmk RB (i.e., rat MuSK-IgG1 receptorbody) was designated "vDmk receptorbody" and deposited with the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland 20852 on May 16, 1995 under ATCC Accession No. VR-2507. The cDNA clone pBL-hAgrin1 was deposited with the American Type Culture Collection, 12301 Parklawn Drive, Rockville, Maryland 20852 on December 12, 1995 under ATCC Accession No. 97378.

15

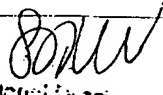
The present invention is not to be limited in scope by the specific embodiments described herein. Indeed, various modifications of the invention in addition to those described herein will become apparent to those skilled in the art from the foregoing description and accompanying figures. Such modifications are intended to fall within the scope of the appended claims.

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INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

(PCT Rule 13bis)

A. The indications made below relate to the microorganism referred to in the description on page <u>86</u> , line <u>5-16</u>	
B. IDENTIFICATION OF DEPOSIT	
Further deposits are identified on an additional sheet <input checked="" type="checkbox"/>	
Name of depositary institution <p style="text-align: center;">American Type Culture Collection</p>	
Address of depositary institution (including postal code and country) <p style="text-align: center;">12301 Parklawn Drive Rockville, Maryland 20852 U.S.A.</p>	
Date of deposit <p style="text-align: center;">July 13, 1993</p>	Accession Number <p style="text-align: center;">75498</p>
C. ADDITIONAL INDICATIONS (leave blank if not applicable) This information is continued on an additional sheet <input type="checkbox"/>	
Applicant wishes that, until publication of the mention of the grant of a European patent or until the date on which the application is refused or withdrawn or is deemed to be withdrawn, the deposit shall be made available as provided in Rule 28(3) of the Implementing Regulations under the European Patent Convention only by the issue of a sample to an expert nominated by the requester (Rule 28(4) of the implementing regulations).	
D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (if the indications are not for all designated States)	
E. SEPARATE FURNISHING OF INDICATIONS (leave blank if not applicable)	
The indications listed below will be submitted to the International Bureau later (specify the general nature of the indications e.g., "Accession Number of Deposit")	

<p>For receiving Office use only</p> <p><input checked="" type="checkbox"/> This sheet was received with the international application</p> <p>Authorized officer:  Susan V. Hall PCT International Bureau</p>	<p>For International Bureau use only</p> <p><input type="checkbox"/> This sheet was received by the International Bureau on:</p> <p>Authorized officer:</p>
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WHAT IS CLAIMED IS:

1. An isolated nucleic acid molecule comprising a nucleotide sequence encoding the active portion of human agrin, wherein the nucleotide sequence is selected from the group consisting of:
 - (a) the nucleotide sequence comprising the coding region of the active portion of human agrin contained in the vector designated as pBL-hAgrin 1 (ATCC Accession No. 97378);
 - (b) a nucleotide sequence that hybridizes under stringent conditions to the nucleotide sequence of (a) and which encodes the active portion of human agrin; and
 - (c) a nucleotide sequence that, as a result of the degeneracy of the genetic code, differs from the nucleotide sequence of (a) or (b) and which encodes the active portion of human agrin.
2. An isolated nucleic acid molecule comprising a nucleotide sequence encoding the active portion of human agrin, wherein the nucleotide sequence is selected from the group consisting of:
 - (a) the nucleotide sequence as set forth in Figure 15;
 - (b) the nucleotide sequence encoding amino acids 24 to 492 as set forth in Figure 15;
 - (c) the nucleotide sequence encoding amino acids 60 to 492 as set forth in Figure 15;
 - (d) the nucleotide sequence encoding amino acids 76 to 492 as set forth in Figure 15;
 - (e) the nucleotide sequence encoding amino acids 126 to 492 as set forth in Figure 15;
 - (f) the nucleotide sequence encoding amino acids 178 to 492 as set forth in Figure 15;
 - (g) the nucleotide sequence encoding amino acids 222 to 492 as set forth in Figure 15;

- (h) the nucleotide sequence encoding amino acids 260 to 492 as set forth in Figure 15;
 - (i) the nucleotide sequence encoding amino acids 300 to 492 as set forth in Figure 15;
 - (j) a nucleotide sequence that hybridizes under stringent conditions to any of the nucleotide sequences of (a) through (i) and which encodes the active portion of human agrin; and
 - (k) a nucleotide sequence that, as a result of the degeneracy of the genetic code, differs from any of the nucleotide sequences of (a) through (j) and which encodes the active portion of human agrin.
- 3. An isolated nucleic acid molecule of claim 1 or 2, which is lacking an insert at position Y.
 - 4. An isolated nucleic acid molecule of claim 1 or 2, which is lacking an insert at position Z.
 - 5. An isolated polypeptide encoded by the nucleic acid molecule of claim 1, 2, 3 or 4.
 - 6. A polypeptide of claim 5, modified by covalent attachment of a polyethylene glycol molecule.
 - 7. A vector which comprises the isolated nucleic acid molecule of claim 1, 2, 3 or 4.
 - 8. An expression vector comprising a nucleic acid molecule of claim 1, 2, 3 or 4 wherein the nucleic acid molecule is operatively linked to an expression control sequence.

9. A host-vector system for the production of a polypeptide having the biological activity of human agrin which comprises the vector of claim 8, in a suitable host cell.
10. The host-vector system of claim 9, wherein the suitable host cell is a bacterial cell, yeast cell, insect cell, or mammalian cell.
11. A method of producing a polypeptide having the biological activity of human agrin which comprises growing cells of the host-vector system of claim 9 or 10, under conditions permitting production of the polypeptide and recovering the polypeptide so produced.
12. A method of promoting the growth, differentiation or survival of a MuSK receptor expressing cell comprising administering to the cell an effective amount of agrin.
13. The method of claim 12, wherein the MuSK receptor expressing cell is a cell which is normally found in muscle, heart, spleen, ovary or retina.
14. The method of claim 12 or 13, wherein the MuSK receptor expressing cell is a cell which has been genetically engineered to express the MuSK receptor.
15. An antibody capable of specifically binding the polypeptide of claim 5 or 6.
16. A monoclonal antibody of claim 15.
17. A polyclonal antibody of claim 15.

18. A method of detecting the presence of human agrin in a sample comprising:
 - a) reacting the sample with an antibody of claim 15, under conditions whereby the antibody binds to human agrin present in the sample; and
 - b) detecting the bound antibody, thereby detecting the presence of human agrin in the sample.
19. The method of claim 18, wherein the antibody is a polyclonal antibody.
20. The method of claim 18, wherein the antibody is a monoclonal antibody.
21. The method of claim 18, wherein the sample is a biological tissue.
22. The method of claim 18, wherein the sample is a body fluid.
23. The method of claim 22, wherein the body fluid is selected from the group consisting of cerebrospinal fluid, blood, serum, plasma, urine and saliva.
24. The method of claim 18, wherein the sample is a cell extract.
25. The host-vector system of claim 10, wherein the bacterial cell is E. coli.
26. The host-vector system of claim 10, wherein the yeast cell is Pichia pastoris.
27. The host-vector system of claim 10, wherein the insect cell is Spodoptera frugiperda.

28. The host-vector system of claim 10, wherein the mammalian cell is a COS cell.
29. The host-vector system of claim 10, wherein the mammalian cell is a CHO cell.
30. A method of treating a patient suffering from a disease or disorder affecting muscle comprising administering to the patient an effective amount of the polypeptide of claim 5 or 6, or a derivative thereof.
31. The method of claim 30, wherein the disease or disorder is muscle atrophy resulting from denervation due to nerve trauma, degenerative, metabolic or inflammatory neuropathy, peripheral neuropathy, or damage to nerves caused by environmental toxins or drugs.
32. The method of claim 30, wherein the disease or disorder is muscle atrophy due to a motor neuronopathy.
33. The method of claim 30, wherein the disease or disorder is muscle atrophy due to chronic disuse.
34. The method of claim 30, wherein the disease or disorder is muscle atrophy due to metabolic stress or nutritional insufficiency.
35. The method of claim 30, wherein the disease or disorder is muscle atrophy due to a muscular dystrophy syndrome.
36. The method of claim 30, wherein the disease or disorder is muscle atrophy due to a congenital myopathy.

37. The method of claim 30, wherein the disease or disorder is an acquired (toxic or inflammatory) myopathy.
38. A polypeptide as defined in claim 5 or 6, for use in a method of treatment of the human or animal body by therapy or in a method of diagnosis.
39. A polypeptide according to claim 38, for use in a method of treatment of the human or animal body of a disease or disorder that affects muscle.
40. A polypeptide according to claim 39, wherein the disease or disorder is muscle atrophy resulting from denervation due to nerve trauma, degenerative, metabolic or inflammatory neuropathy, peripheral neuropathy, or damage to nerves caused by environmental toxins or drugs.
41. A polypeptide according to claim 39, wherein the disease or disorder is muscle atrophy due to a motor neuronopathy.
42. A polypeptide according to claim 39, wherein the disease or disorder is muscle atrophy due to chronic disuse.
43. A polypeptide according to claim 39, wherein the disease or disorder is muscle atrophy due to metabolic stress or nutritional insufficiency.
44. A polypeptide according to claim 39, wherein the disease or disorder is muscle atrophy due to a muscular dystrophy syndrome.

45. A polypeptide according to claim 39, wherein the disease or disorder is muscle atrophy due to a congenital myopathy.
46. A polypeptide according to claim 39, wherein the disease or disorder is an acquired (toxic or inflammatory) myopathy.
47. Use of a polypeptide as defined in claim 5 or 6 in the manufacture of a medicament for the treatment of a disease or disorder affecting muscle.
48. A pharmaceutical composition comprising a polypeptide as defined in claim 5 or 6 and a pharmaceutically acceptable carrier.
49. A diagnostic test kit for detecting the presence of human agrin in a sample, said kit comprising an antibody as defined in any of claims 15 to 17, and means for determining whether or not the antibody binds to human agrin, thereby allowing detection of the presence of human agrin in the sample.
50. A method of treating a patient suffering from a disease or disorder affecting muscle comprising administering to the patient an effective amount of the nucleic acid molecule of claim 1, 2, 3 or 4, or a derivative thereof.
51. A nucleic acid molecule as defined in claim 1, 2, 3 or 4, or a derivative thereof, for use in a method of treatment of the human or animal body by therapy or in a method of diagnosis.
52. A nucleic acid molecule according to claim 51, for use in a method of treatment of the human or animal body of a disease or disorder that affects muscle.

53. Use of a nucleic acid molecule as defined in claim 1, 2, 3 or 4, or a derivative thereof, in the manufacture of a medicament for the treatment of a disease or disorder affecting muscle.
54. A pharmaceutical composition comprising a nucleic acid molecule as defined in claim 1, 2, 3 or 4, or a derivative thereof, and a pharmaceutically acceptable carrier.
55. A nucleic acid molecule according to claim 1, 2, 3 or 4, or a derivative thereof, substantially as hereinbefore described.
56. A vector according to claim 7 or an expression vector according to claim 8, substantially as hereinbefore described.
57. A host vector system according to claim 9, substantially as hereinbefore described.
58. A method according to claim 11, 12, 18, 30 or 50 substantially as hereinbefore described.
59. An antibody according to claim 15 substantially as hereinbefore described.
60. A polypeptide according to claim 39 substantially as hereinbefore described.
61. Use according to claim 47 or 53 substantially as hereinbefore described.
62. A pharmaceutical composition according to claim 48 or 54 substantially as hereinbefore described.

63. A kit according to claim 49 substantially as hereinbefore described.

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Fig.1.

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      5   10   15   20   25   30   35   40   45   50   55   60   65   70
      *   *   *   *   *   *   *   *   *   *   *   *   *
GAATTGGCA OGAGCAACA GTCATTAGTG GAAGACTCTA TTGTAATAAA CTGTGCTTAA AAATGTAAC

      75   80   85   90   95  100  105  110  115  120  125  130  135
      *   *   *   *   *   *   *   *   *   *   *   *
CAGGAGGGT TTTTTTCTC CACATGTGC AGAAGCAAC TTCTTCTCG AGCTGGATT AATC ATG
      HD>

140   145   150   155   160   165   170   175   180   185   190
      *   *   *   *   *   *   *   *   *   *   *
AGA GAG CTC GTC AAC ATT CCA CTG TTA CAG ATG CTC ACC CTG GTT GGC TTC AGC GGG
R   E   L   V   N   I   P   L   L   Q   H   L   T   L   V   A   F   S   G>

195   200   205   210   215   220   225   230   235   240   245   250
      *   *   *   *   *   *   *   *   *   *   *
ACC GAG AAA CTT CCA AAA GGC OCT GTC ATC ACC ACG OCT CTT GAA ACT GTA GAT GGC
T   E   K   L   P   K   A   P   V   I   T   T   P   L   E   T   V   D   A>

255   260   265   270   275   280   285   290   295   300   305
      *   *   *   *   *   *   *   *   *   *   *
TTA GTT GAA GAA GTG GCG ACT TTC ATG TGC GGC GTG GAA TOC TAC OCT CAG OCT GAA
L   V   E   E   V   A   T   F   H   C   A   V   E   S   Y   P   Q   P   E>

310   315   320   325   330   335   340   345   350   355   360   365
      *   *   *   *   *   *   *   *   *   *   *
ATT TCT TGG ACC AGA AAT AAA ATT CTC ATC AAG CTG TTT GAC ACC GGC TAC AGC ATC
I   S   W   T   R   N   K   I   L   I   K   L   F   D   T   R   Y   S   I>

370   375   380   385   390   395   400   405   410   415   420
      *   *   *   *   *   *   *   *   *   *   *
CGA GAG AAC GGT CAG CTC CTC ACC ATC CTG AGT GTG GAG GAC AGT GAT GAT GGC ATC
R   E   N   G   Q   L   L   T   I   L   S   V   E   D   S   D   D   G   I>

425   430   435   440   445   450   455   460   465   470   475
      *   *   *   *   *   *   *   *   *   *   *
TAC TGC TGC ACA GGC AAC AAT GGA GTG GGA GGA GCG GTG GAA AGT TGT GGC GGC CTG
Y   C   C   T   A   N   N   G   V   G   G   A   V   E   S   C   G   A   L>

480   485   490   495   500   505   510   515   520   525   530   535
      *   *   *   *   *   *   *   *   *   *   *
CAA GTG AAG ATG AAG OCT AAA ATA ACT OGT OCT OCC ATC AAT GTA AAA ATA ATT GAG
Q   V   K   H   K   P   K   I   T   R   P   P   I   N   V   K   I   I   E>

540   545   550   555   560   565   570   575   580   585   590
      *   *   *   *   *   *   *   *   *   *   *
GGA TTG AAA GCA GTC CTA CCG TGC ACT ACG ATG GGT AAC OCC AAG CCA TOC GTG TOC
G   L   K   A   V   L   P   C   T   T   H   G   N   P   K   P   S   V   S>

595   600   605   610   615   620   625   630   635   640   645   650
      *   *   *   *   *   *   *   *   *   *   *
TGG ATT AAG GGG GAC AGT GCT CTC AGG GAA AAT TOC AGG ATT GCA GTT CTT GAA TCT
W   I   K   G   D   S   A   L   R   E   N   S   R   I   A   V   L   E   S>

655   660   665   670   675   680   685   690   695   700   705
      *   *   *   *   *   *   *   *   *   *   *
GGG AGT TTA AGG ATC CAT AAT GTG CAA AAG GAA GAC GCA GGA CAG TAC CGA TGT GTG
G   S   L   R   I   H   N   V   Q   K   E   D   A   G   Q   Y   R   C   V>

710   715   720   725   730   735   740   745   750   755   760
      *   *   *   *   *   *   *   *   *   *   *
GCA AAA AAC AGC CTG GGC ACA GCT TAC TOC AAA CTG GTG AAG CTG GAA GTG GAG GTT
A   K   N   S   L   G   T   A   Y   S   K   L   V   K   L   E   V   E   V>

765   770   775   780   785   790   795   800   805   810   815   820
      *   *   *   *   *   *   *   *   *   *   *
TTT GCA AGA ATC CTG CGT GCT OCT GAA TOC CAC AAT GTC ACC TTT GGT TOC TTT GTA
F   A   R   I   L   R   A   P   E   S   H   N   V   T   F   G   S   F   V>

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Fig.1 (Cont 1).

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      825   830   835   840   845   850   855   860   865   870   875
      *   *   *   *   *   *   *   *   *   *
AOC CTA CGC TGC ACA GCA ATA GGC ATG OCT GTC COC ACC ATC AGC TGG ATT GAA AAC
T   L   R   C   T   A   I   G   M   P   V   P   T   I   S   W   I   E   N>

880   885   890   895   900   905   910   915   920   925   930   935
      *   *   *   *   *   *   *   *   *   *
GGA AAT GCT GTT TCT TCA GGT TOC ATT CAA GAG AAT GTG AAA GAC OGA GTG ATT GAC
G   N   A   V   S   S   G   S   I   Q   E   N   V   K   D   R   V   I   D>

      940   945   950   955   960   965   970   975   980   985   990
      *   *   *   *   *   *   *   *   *   *
TCA AGA CTC CAG CTC TTT ATC ACA AAG OCA GGA CTC TAC ACA TGC ATA GCT ACC AAT
S   R   L   Q   L   F   I   T   K   P   G   L   Y   T   C   I   A   T   N>

995   1000   1005   1010   1015   1020   1025   1030   1035   1040   1045
      *   *   *   *   *   *   *   *   *   *
AAG CAT GGA GAG AAA TTC AGT AOC GCA AAG GCT GCA GOC ACT GTC AGT ATA GCA GAA
K   H   G   E   K   F   S   T   A   K   A   A   A   T   V   S   I   A   E>

1050  1055   1060   1065   1070   1075   1080   1085   1090   1095   1100   1105
      *   *   *   *   *   *   *   *   *   *
TGG AGC AAA TCA CAG AAA GAA AGC AAA GGC TAC TGT GOC CAG TAC AGA GGG GAG GTG
W   S   K   S   Q   K   E   S   K   G   Y   C   A   Q   Y   R   G   E   V>

      1110  1115   1120   1125   1130   1135   1140   1145   1150   1155   1160
      *   *   *   *   *   *   *   *   *   *
TGT GAT GGC GTC CTG GTG AAA GAC TCT CTT GTC TTC TTC AAC AOC TOC TAT OCC GAC
C   D   A   V   L   V   K   D   S   L   V   F   F   N   T   S   Y   P   D>

1165  1170  1175   1180   1185   1190   1195   1200   1205   1210   1215   1220
      *   *   *   *   *   *   *   *   *   *
OCT GAG GAG GOC CAA GAG CTG CTG ATC CAC ACT GOG TGG AAT GAA CTC AAG GCT GTG
P   E   E   A   Q   E   L   L   I   H   T   A   W   N   E   L   K   A   V>

      1225  1230  1235   1240   1245   1250   1255   1260   1265   1270   1275
      *   *   *   *   *   *   *   *   *   *
AGC OCA CTC TGC OGA OCA GCT GCC GAG GCT CTG CTG TGT AAT CAC CTC TTC CAG GAG
S   P   L   C   R   P   A   A   E   A   L   L   C   N   H   L   F   Q   E>

1280  1285   1290  1295   1300   1305   1310   1315   1320  1325   1330
      *   *   *   *   *   *   *   *   *   *
TGC AGC OCT GGA GTG CTA OCT ACT OCT ATG COC ATT TGC AGA GAG TAC TGC TTG GCA
C   S   P   G   V   L   P   T   P   M   P   I   C   R   E   Y   C   L   A>

1335  1340  1345   1350  1355   1360   1365  1370   1375   1380  1385   1390
      *   *   *   *   *   *   *   *   *   *
GTA AAG GAG CTC TTC TGT GCA AAG GAA TGG CTG GCA ATG GAA GGG AAG ACC CAC CGC
V   K   E   L   F   C   A   K   E   W   L   A   M   E   G   K   T   H   R>

      1395  1400   1405   1410  1415   1420   1425  1430   1435   1440  1445
      *   *   *   *   *   *   *   *   *   *
GGA CTC TAC AGA TOC GGG ATG CAT TTC CTC COG GTC COG GAG TGC AGC AAG CTT OCC
G   L   Y   R   S   G   M   H   F   L   P   V   P   E   C   S   K   L   P>

1450  1455  1460   1465   1470  1475   1480   1485  1490   1495   1500  1505
      *   *   *   *   *   *   *   *   *   *
AGC ATG CAC CAG GAC CCC ACA GCC TGC ACA AGA-CTG COG TAT TTA GAT TAT AAA AAA
S   H   H   Q   D   P   T   A   C   T   R   L   P   Y   L   D   Y   K   K>

      1510  1515  1520   1525   1530  1535   1540   1545  1550   1555   1560
      *   *   *   *   *   *   *   *   *   *
GAA AAC ATA ACA ACA TTC CCG TOC ATA ACG TOC TOC AAG COG AGC GTG GAC ATT OCA
E   N   I   T   T   F   P   S   I   T   S   S   K   P   S   V   D   I   P>

1565  1570   1575  1580   1585   1590  1595   1600   1605  1610   1615
      *   *   *   *   *   *   *   *   *   *
AAC CTG OCT GCC TOC ACC TCT TOC TTC GOC GTC TOG CCT GOG TAC TOC ATG ACT GTC
N   L   P   A   S   T   S   S   F   A   V   S   P   A   Y   S   H   T   V>

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Fig.1 (Cont 2).

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1620 1625 1630 1635 1640 1645 1650 1655 1660 1665 1670 1675
  *   *   *   *   *   *   *   *   *   *   *
ATC ATC TOC ATC ATG TOC TGC TTT GCG GTG TTT GCT CTC CTC ACC ATC ACT ACT CTC
I  I  S  I  M  S  C  F  A  V  F  A  L  L  T  I  T  T  L>

1680 1685 1690 1695 1700 1705 1710 1715 1720 1725 1730
  *   *   *   *   *   *   *   *   *   *   *
TAT TGC TGC CGA AGG AGG AGA GAG TGG AAA AAT AAG AAA AGA GAG TOG GCA GCG GTG
Y  C  C  R  R  R  R  E  W  K  N  K  K  R  E  S  A  A  V>

1735 1740 1745 1750 1755 1760 1765 1770 1775 1780 1785 1790
  *   *   *   *   *   *   *   *   *   *   *
ACC CTC ACC ACA TTG OCT TOC GAG CTC CTG CTG GAC AGG CTG CAT CCC AAC CCC ATG
T  L  T  T  L  P  S  E  L  L  L  D  R  L  H  P  N  P  M>

1795 1800 1805 1810 1815 1820 1825 1830 1835 1840 1845
  *   *   *   *   *   *   *   *   *   *   *
TAC CAG AGG ATG CCA CTC CTT CTG AAT CCC AAG TTG CTC AGC CTG GAG TAT CCG AGG
Y  Q  R  M  P  L  L  L  N  P  K  L  L  S  L  E  Y  P  R>

1850 1855 1860 1865 1870 1875 1880 1885 1890 1895 1900
  *   *   *   *   *   *   *   *   *   *   *
AAT AAC ATC GAG TAT GTC AGA GAC ATC GGA GAG GGA GCG TTT GGA AGG GTC TTT CAA
N  N  I  E  Y  V  R  D  I  G  E  G  A  F  G  R  V  F  Q>

1905 1910 1915 1920 1925 1930 1935 1940 1945 1950 1955 1960
  *   *   *   *   *   *   *   *   *   *   *
GCG AGG GCG CCA GGC TTG CTT OCT TAT GAA CCC TTC ACT ATG GTG GCT GTG AAG ATG
A  R  A  P  G  L  L  P  Y  E  P  F  T  M  V  A  V  K  M>

1965 1970 1975 1980 1985 1990 1995 2000 2005 2010 2015
  *   *   *   *   *   *   *   *   *   *   *
CTG AAG GAG GAG GCG TOC GCA GAT ATG CAG GCA GAC TTT CAG AGG GAG GCA GCG CTC
L  K  E  E  A  S  A  D  M  Q  A  D  F  Q  R  E  A  A  L>

2020 2025 2030 2035 2040 2045 2050 2055 2060 2065 2070 2075
  *   *   *   *   *   *   *   *   *   *   *
ATG GCG GAG TTT GAC AAC CCC AAC ATT GTG AAG CTC TTA GGT GTG TGT GCT GTT GGG
M  A  E  F  D  N  P  N  I  V  K  L  L  G  V  C  A  V  G>

2080 2085 2090 2095 2100 2105 2110 2115 2120 2125 2130
  *   *   *   *   *   *   *   *   *   *   *
AAG CCA ATG TGC CTG CTC TTT GAA TAT ATG GCG TAT GGT GAC CTC AAT GAG TTC CTC
K  P  M  C  L  L  F  E  Y  M  A  Y  G  D  L  N  E  F  L>

2135 2140 2145 2150 2155 2160 2165 2170 2175 2180 2185
  *   *   *   *   *   *   *   *   *   *   *
CGA AGC ATG TOC OCT CAC ACT GTG TGC AGC CTC AGC CAC AGT GAC CTG TOC ACG AGG
R  S  M  S  P  H  T  V  C  S  L  S  H  S  D  L  S  T  R>

2190 2195 2200 2205 2210 2215 2220 2225 2230 2235 2240 2245
  *   *   *   *   *   *   *   *   *   *   *
GCT GCG GTG TOC AGC OCT GGT OCT CCA CCC CTG TCT TGT GCG GAA CAG CTC TGT ATT
A  R  V  S  S  P  G  P  P  P  L  S  C  A  E  Q  L  C  I>

2250 2255 2260 2265 2270 2275 2280 2285 2290 2295 2300
  *   *   *   *   *   *   *   *   *   *   *
GCG AGG CAA GTG GCA GCT GCG ATG GCC TAC CTG TOG GAG CGC AAG TTT GTC CAT GCG
A  R  Q  V  A  A  G  M  A  Y  L  S  E  R  K  F  V  H  R>

2305 2310 2315 2320 2325 2330 2335 2340 2345 2350 2355 2360
  *   *   *   *   *   *   *   *   *   *   *
GAC TTA GCT ACC AGG AAC TGC CTG GTT GGA GAG AAC ATG GTG GTG AAA ATT GCA GAC
D  L  A  T  R  N  C  L  V  G  E  N  M  V  V  K  I  A  D>

2365 2370 2375 2380 2385 2390 2395 2400 2405 2410 2415
  *   *   *   *   *   *   *   *   *   *   *
TTT GCG CTC TCT AGG AAC ATC TAC TCC CCA GAC TAC TAC AAA GCT GAT GGA AAC GAT
F  G  L  S  R  N  I  Y  S  A  D  Y  Y  K  A  D  G  N  D>

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Fig.1 (Cont 3).

```

2420 2425 2430 2435 2440 2445 2450 2455 2460 2465 2470
*      *      *      *      *      *      *      *      *      *
GCT ATA OCT ATC OGC TGG ATG OCA OCC GAG TCT ATC TTC TAC AAC OGC TAC AOC ACG
A I P I R W M P P E S I F Y N R Y T T>

2475 2480 2485 2490 2495 2500 2505 2510 2515 2520 2525 2530
*      *      *      *      *      *      *      *      *      *
GAG TCA GAT GTG TGG OCT TAT GGC GTG GTC CTC TGG GAG ATC TTC TOC TAT GGA CTG
E S D V W A Y G V V L W E I F S Y G L>

2535 2540 2545 2550 2555 2560 2565 2570 2575 2580 2585
*      *      *      *      *      *      *      *      *      *
CAG OCC TAC TAT GGA ATG GOC CAT GAG GAG GTC ATT TAC TAT GTG AGA GAT GGT AAC
Q P Y Y G H A H E E V I Y Y V R D G RD>

2590 2595 2600 2605 2610 2615 2620 2625 2630 2635 2640 2645
*      *      *      *      *      *      *      *      *      *
ATC CTT GOC TGC OCT GAG AAC TGT OCC TTG GAA CTG TAC AAC CTT ATG OGC CTA TGT
I L A C P E H C P L E L Y N L H R L C>

2650 2655 2660 2665 2670 2675 2680 2685 2690 2695 2700
*      *      *      *      *      *      *      *      *      *
TGG AGC AAG CTG OCT GCA GAC AGA OCC AGC TTC TGC AGT ATC CAC OGC ATC CTG CAG
W S K L P A D R P S F C S I H R I L Q>

2705 2710 2715 2720 2725 2730 2735 2740 2745 2750 2755 2760
*      *      *      *      *      *      *      *      *      *
OGC ATG TGC GAG AGA GCA GAG GGA ACG GTA GGC GTC TAA GGTGACCA TGCTCAACA
R M C E R A E G T V G V *>

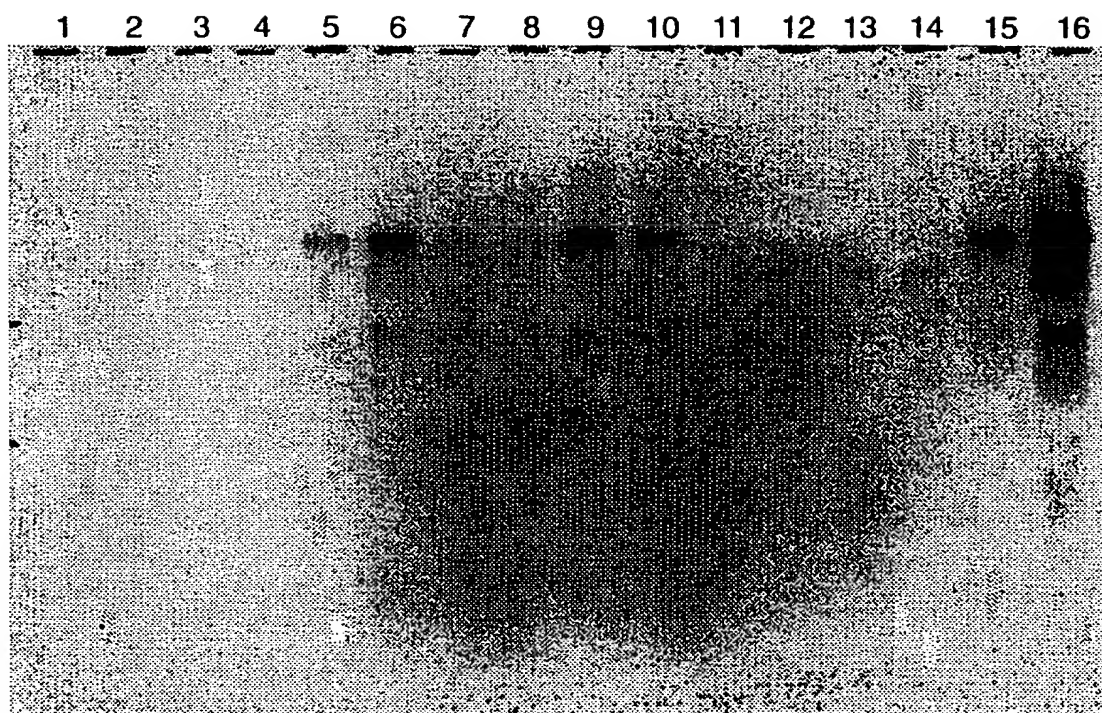
2765 2770 2775 2780 2785 2790 2795 2800 2805 2810 2815 2820 2825 2830
*      *      *      *      *      *      *      *      *      *
ACAOCAGGA GGATCTTTTC AGACTGOGAG CTGGAGGGAT OCTAAAGCAG AGGGGQATA AGTNCAGATA

2835 2840 2845 2850 2855 2860 2865
*      *      *      *      *
GGAAGAGTTT ATCTCAGGCA GCAOGTNCAG TTGGTTGTT

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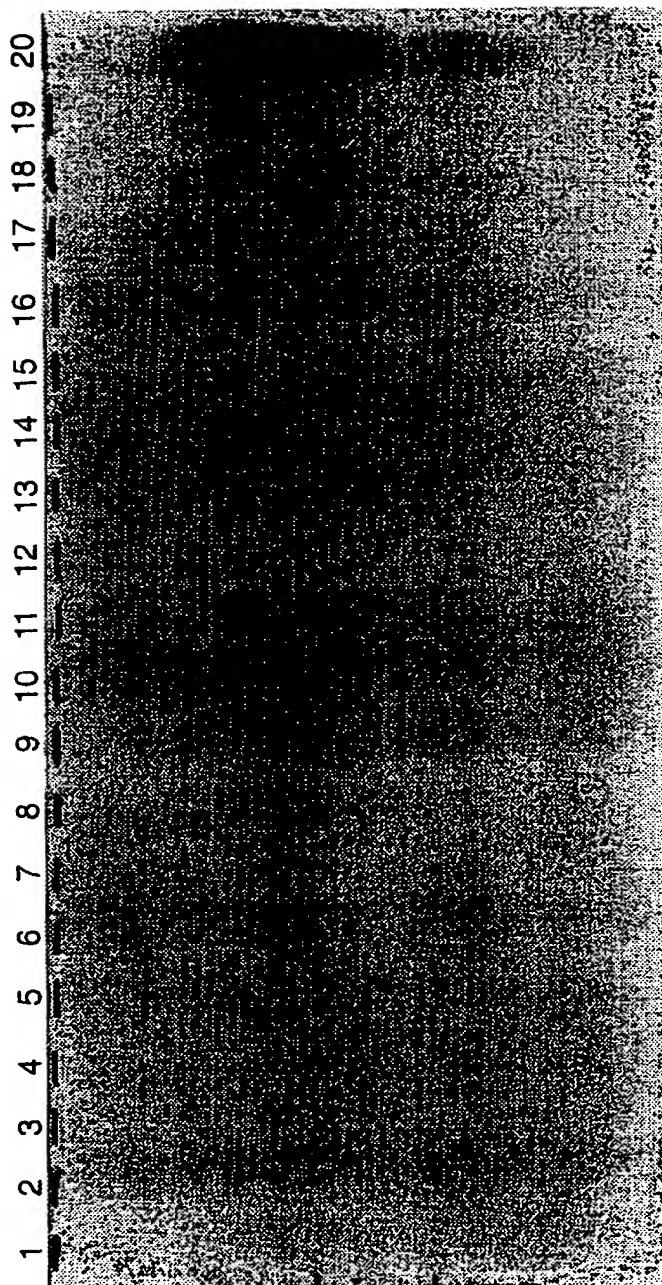
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Fig.2.



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Fig.3.



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Fig.4.

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      10      20      30      40      50      60
      .      .      .      .      .      .
ATG AGA GAG CTC GTC AAC ATT OCA CTG GTA CAT ATT CTT ACT CTG GTT GGC TTC AOC GGA
H R E L V N I P L V H I L T L V A F S Q

      70      80      90      100      110      120
      .      .      .      .      .      .
ACT GAG AAA CTT OCA AAA GCT OCT GTC ATC AOC ACT OCT CTT GAA ACA GTG GAT GGC TTA
T E K L P K A P V I T T P L E T V D A L

      130      140      150      160      170      180
      .      .      .      .      .      .
GTT GAA GAA GTG OCT ACT TTC ATG TGT GCA GTG GAA TCC TAC OOC CAG OCT GAG ATT TCC
V E E V A T F H C A V E S Y P Q P E I S

      190      200      210      220      230      240
      .      .      .      .      .      .
TGG ACT AGA AAT AAA ATT CTC ATT AAA CTC TTT GAC AOC OOC TAC AOC ATC OOG GAG AAT
W T R N K I L I K L F D T R Y S I R E N

      250      260      270      280      290      300
      .      .      .      .      .      .
GGG CAG CTC CTC ACC ATC CTG AGT GTG GAA GAC AGT GAT GAT GGC ATT TAC TGC TCC ACG
G Q L L T I L S V E D S D D G I Y C C T

      310      320      330      340      350      360
      .      .      .      .      .      .
GCC AAC AAT GGT GTG GGA GGA GCT GTG GAG AGT TGT GGA GGC CTG CAA GTG AAG ATG AAA
A N N G V G G A V E S C G A L Q V K H K

      370      380      390      400      410      420
      .      .      .      .      .      .
OCT AAA ATA ACT CGC CCT CCC ATA AAT GTG AAA ATA ATA GAG GGA TTA AAA GCA GTC CTA
P K I T R P P I N V K I I E G L K A V L

      430      440      450      460      470      480
      .      .      .      .      .      .
CCA TGT ACT ACA ATG GGT AAT CCC AAA CCA TCA GTG TCT TGG ATA AAG GGA GAC AGC CCT
P C T T H G N P K P S V S W I K G D S P

      490      500      510      520      530      540
      .      .      .      .      .      .
CTC AGG GAA AAT TCC CGA ATT GCA GTT CTT GAA TCT GGG AGC TTG AGG ATT CAT AAC GTA
L R E N S R I A V L E S G S L R I H N V

      550      560      570      580      590      600
      .      .      .      .      .      .
CAA AAG GAA GAT GCA GGA CAG TAT CGA TGT GTG GCA AAA AAC AGC CTC GGG ACA GCA TAT
Q K E D A G Q Y R C V A K N S L G T A Y

      610      620      630      640      650      660
      .      .      .      .      .      .
TCC AAA GTG GTG AAG CTG GAA GTT GAG GTT TTT GCC AGG ATC CTG CGG GCT CCT GAA TCC
S K V V K L E V E V F A R I L R A P E S

      670      680      690      700      710      720
      .      .      .      .      .      .
CAC AAT GTC ACC TTT GGC TCC TTT GTG ACC CTG CAC TGT ACA GCA ACA GGC ATT CCT GTC
H N V T F G S F V T L H C T A T G I P V

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Fig.4 (Cont 1).

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      730      740      750      760      770      780
      .       .       .       .       .       .
OCC ACC ATC ACC TGG ATT GAA AAC GGA AAT OCT GTT TCT TCT GGG TCC ATT CAA GAG AGT
P   T   I   T   W   I   E   N   G   N   A   V   S   S   G   S   I   Q   E   S

      790      800      810      820      830      840
      .       .       .       .       .       .
GTG AAA GAC CGA GTG ATT GAC TCA AGA CTG CAG CTG TTT ATC ACC AAG CCA GGA CTC TAC
V   K   D   R   V   I   D   S   R   L   Q   L   F   I   T   K   P   G   L   Y

      850      860      870      880      890      900
      .       .       .       .       .       .
ACA TCC ATA GCT ACC AAT AAG CAT GGG GAG AAG TTC AGT ACT GCC AAG GCT GCA GCC ACC
T   C   I   A   T   N   K   H   G   E   K   F   S   T   A   K   A   A   A   T

      910      920      930      940      950      960
      .       .       .       .       .       .
ATC AGC ATA GCA GAA TGG AGT AAA CCA CAG AAA GAT AAC AAA GGC TAC TCC GCC CAG TAC
I   S   I   A   E   W   S   K   P   Q   K   D   N   K   G   Y   C   A   Q   Y

      970      980      990      1000      1010      1020
      .       .       .       .       .       .
AGA GGG GAG GTG TGT AAT GCA GTC CTG GCA AAA GAT OCT CTT GTT TTT CTC AAC ACC TCC
R   G   E   V   C   N   A   V   L   A   K   D   A   L   V   F   L   N   T   S

      1030      1040      1050      1060      1070      1080
      .       .       .       .       .       .
TAT GCG GAC CCT GAG GAG GCC CAA GAG CTA CTG GTC CAC ACG GCC TGG AAT GAA CTG AAA
Y   A   D   P   E   E   A   Q   E   L   L   V   H   T   A   W   N   E   L   K

      1090      1100      1110      1120      1130      1140
      .       .       .       .       .       .
GTA GTG AGC CCA GTC TGC CCG CCA GCT GCT GAG GCT TTG TTG TGT AAC CAC ATC TTC CAG
V   V   S   P   V   C   R   P   A   A   E   A   L   L   C   N   H   I   F   Q

      1150      1160      1170      1180      1190      1200
      .       .       .       .       .       .
GAG TGC AGT CCT GGA GTA GTG CCT ACT CCT ATT CCC ATT TGC AGA GAG TAC TGC TTG GCA
E   C   S   P   G   V   V   P   T   P   I   P   I   C   R   E   Y   C   L   A

      1210      1220      1230      1240      1250      1260
      .       .       .       .       .       .
GTA AAG GAG CTC TTC TGC GCA AAA GAA TGG CTG GTA ATG GAA GAG AAG ACC CAC AGA GGA
V   K   E   L   F   C   A   K   E   W   L   V   H   E   E   K   T   H   R   G

      1270      1280      1290      1300      1310      1320
      .       .       .       .       .       .
CTC TAC AGA TCC GAG ATG CAT TTG CTG TCC GTG CCA GAA TGC AGC AAG CTT CCC AGC ATG
L   Y   R   S   E   H   H   L   L   S   V   P   E   C   S   K   L   P   S   H

      1330      1340      1350      1360      1370      1380
      .       .       .       .       .       .
CAT TGG GAC CCC ACG GCC TGT GCC AGA CTG CCA CAT CTA GAT TAT AAC AAA GAA AAC CTA
H   W   D   P   T   A   C   A   R   L   P   H   L   D   Y   N   K   E   N   L

      1390      1400      1410      1420      1430      1440
      .       .       .       .       .       .
AAA ACA TTC CCA CCA ATG ACG TCC TCA AAG CCA AGT GTG GAC ATT CCA AAT CTG CCT TCC
K   T   F   P   P   H   T   S   S   K   P   S   V   D   I   P   N   L   P   S

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Fig.4 (Cont 2).

1450	1460	1470	1480	1490	1500
TCC	TCC	TCT	TCT	TCC	TTC
S	S	S	S	S	F
TCT	TCT	TCC	TTC	TCT	GTC
S	S	S	F	S	V
TCA	OCT	ACA	TAC	TCC	ATG
S	P	T	Y	S	H
ACT	GTA	ATA	ATC	TCC	ATC
T	V	I	I	S	I
1510	1520	1530	1540	1550	1560
ATG	TCC	AGC	TTT	GCA	ATA
H	S	S	F	A	I
TTT	GTC	CTT	CTT	ACC	ATA
F	V	L	L	T	I
ACT	ACT	CTC	TAT	TGC	TGC
T	T	L	Y	C	C
CGA	AGA				
R	R				
1570	1580	1590	1600	1610	1620
AGA	AAA	CAA	TGG	AAA	AAT
R	K	Q	W	K	N
AAG	AAA	AGA	GAA	TCA	GCA
K	K	R	E	S	A
GTA	AOC	CTC	ACC	ACA	CTG
V	T	L	T	T	L
P					
1630	1640	1650	1660	1670	1680
TCT	GAG	CTC	TTA	CTA	GAT
S	E	L	L	L	D
AGA	CTT	CAT	CCC	AAC	CCC
R	L	H	P	N	P
ATG	TAC	CAG	AGG	ATG	CCG
H	Y	Q	R	H	P
CTC	CTT				
L	L				
1690	1700	1710	1720	1730	1740
CTG	AAC	CCC	AAA	TTG	CTC
L	N	P	K	L	L
AGC	CTG	GAG	TAT	CCA	AGG
S	L	E	Y	P	R
AAT	AAC	ATT	GAA	TAT	GTG
N	N	I	E	Y	V
AGA	GAC				
R	D				
1750	1760	1770	1780	1790	1800
ATC	GGA	GAG	GGA	GCG	TTT
I	G	E	G	A	F
GGA	AGG	GTG	TTT	CAA	GCA
R	V	F	Q	A	R
OCA	OCA	OCA	GCC	TTA	CTT
A	P	G	L	L	P
TAT					
Y					
1810	1820	1830	1840	1850	1860
GAA	CCT	TTC	ACT	ATG	GTG
E	P	F	T	H	V
GCA	GTA	AAG	ATG	CTC	AAA
A	V	K	H	L	K
GAA	GAA	GCC	TGG	GCA	GAT
E	E	A	S	A	D
ATG	CAA				
H	Q				
1870	1880	1890	1900	1910	1920
GCG	GAC	TTT	CAG	AGG	GAG
A	D	F	Q	R	E
GCA	GCC	CTC	ATG	GCA	GAA
A	A	L	H	A	E
TTT	GAC	AAC	OCT	AAC	ATT
F	D	N	P	N	I
GTG	AAG				
V	K				
1930	1940	1950	1960	1970	1980
CTA	TTA	GGA	GTG	TGT	GCT
L	L	G	V	C	A
GTC	GGG	AAG	CCA	ATG	TGC
G	K	P	H	C	L
CTG	CTC	TTT	GAA	TAC	ATG
L	L	F	E	Y	H
GCC	TAT				
A	Y				
1990	2000	2010	2020	2030	2040
GGT	GAC	CTC	AAT	GAG	TTC
G	D	L	N	E	F
CTC	CGC	AGC	ATG	TCC	OCT
L	R	S	H	S	P
CAC	ACC	GTG	TGC	AGC	CTC
H	T	V	C	S	L
AGT	CAC				
S	H				
2050	2060	2070	2080	2090	2100
AGT	GAC	TTG	TCT	ATG	AGG
S	D	L	S	H	R
GCT	CAG	GTC	TCC	AGC	OCT
A	Q	V	S	S	P
GGG	CCC	CCA	CCC	CTC	TCC
G	P	P	P	L	S
TGT	GCT				
C	A				
2110	2120	2130	2140	2150	2160
GAG	CAG	CTT	TGC	ATT	GCC
E	Q	L	C	I	A
AGG	CAG	GTG	GCA	GCT	GGC
R	Q	V	A	A	G
ATG	GCT	TAC	CTC	TCA	GAA
H	A	Y	L	S	E
CGT	AAG				
R	K				

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Fig.4 (Cont 3).

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      2170      2180      2190      2200      2210      2220
      .        .        .        .        .        .
TTT GTT CAC CGA GAT TTA GCC ACC AGG AAC TGC CTG GTG GGC GAG AAC ATG GTG GTG AAA
F   V   H   R   D   L   A   T   R   N   C   L   V   G   E   N   H   V   V   K

      2230      2240      2250      2260      2270      2280
      .        .        .        .        .        .
ATT GCC GAC TTT GGC CTC TCC AGG AAC ATC TAC TCA OCA GAC TAC TAC AAA GCT AAT GAA
I   A   D   F   G   L   S   R   N   I   Y   S   A   D   Y   Y   K   A   N   E

      2290      2300      2310      2320      2330      2340
      .        .        .        .        .        .
AAC GAC GCT ATC OCT ATC CGT TGG ATG OCA OCA GAG TCC ATT TTT TAT AAC CGC TAC ACT
N   D   A   I   P   I   R   W   H   P   P   E   S   I   F   Y   N   R   Y   T

      2350      2360      2370      2380      2390      2400
      .        .        .        .        .        .
ACA GAG TCT GAT GTG TGG GCC TAT GGC GTG GTC CTC TGG GAG ATC TTC TCC TAT GGC CTG
T   E   S   D   V   W   A   Y   G   V   V   L   W   E   I   F   S   Y   G   L

      2410      2420      2430      2440      2450      2460
      .        .        .        .        .        .
CAG CCC TAC TAT GGG ATG GCC CAT GAG GAG GTC ATT TAC TAC GTG CGA GAT GGC AAC ATC
Q   P   Y   Y   G   H   A   H   E   E   V   I   Y   Y   V   R   D   G   N   I

      2470      2480      2490      2500      2510      2520
      .        .        .        .        .        .
CTC TCC TGC OCT GAG AAC TGC CCC GTG GAG CTG TAC AAT CTC ATG CGT CTA TGT TGG AGC
L   S   C   P   E   N   C   P   V   E   L   Y   N   L   H   R   L   C   W   S

      2530      2540      2550      2560      2570      2580
      .        .        .        .        .        .
AAG CTG OCT GCA GAC AGA CCC AGT TTC ACC AGT ATT CAC CGA ATT CTG GAA CGC ATG TGT
K   L   P   A   D   R   P   S   F   T   S   I   H   R   I   L   E   R   H   C

      2590      2600      2610
      .        .        .
GAG AGG GCA GAG GGA ACT GTG AGT GTC TAA
E   R   A   E   G   T   V   S   V   *

```


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Fig.5.

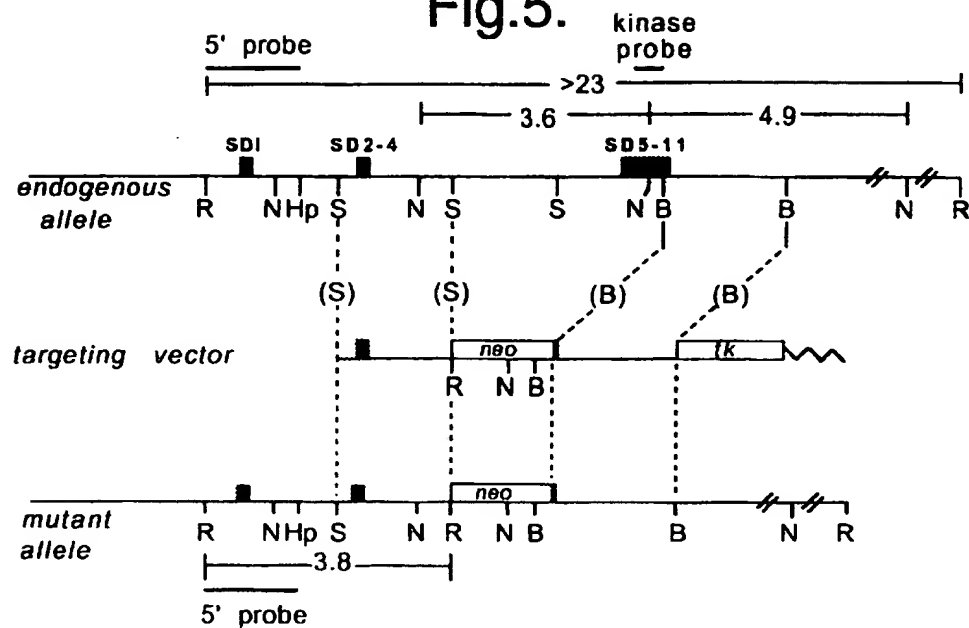
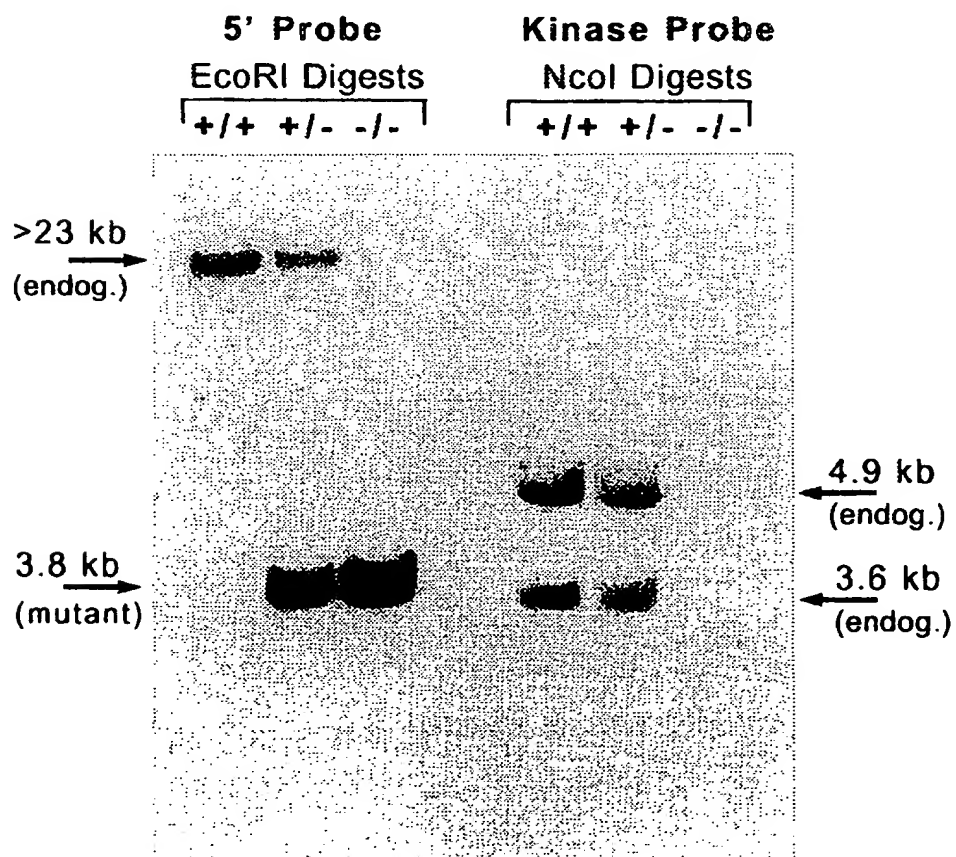


Fig.6.



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Fig.7B.



Fig.7A.



Fig.7D.

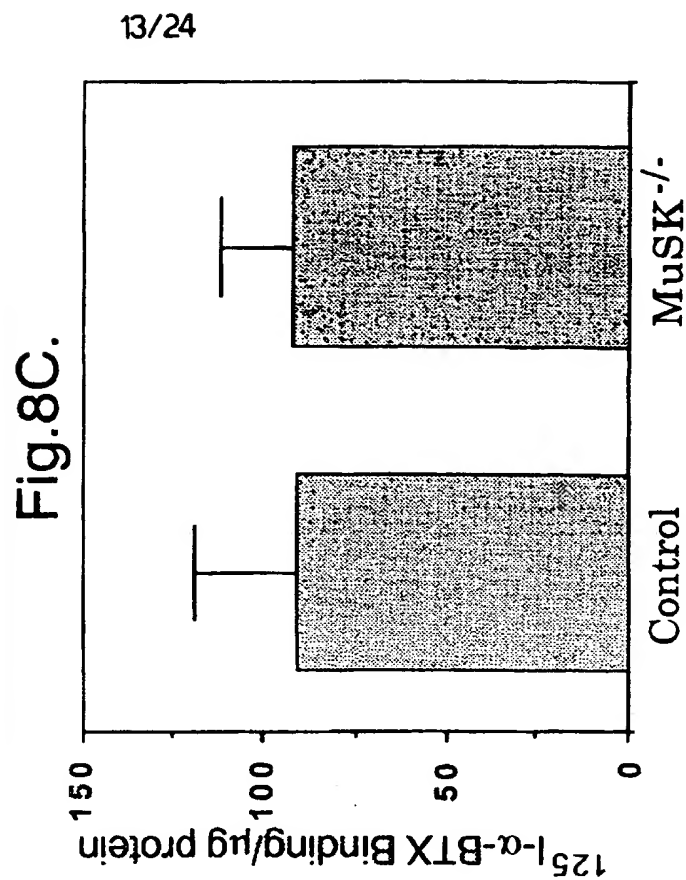
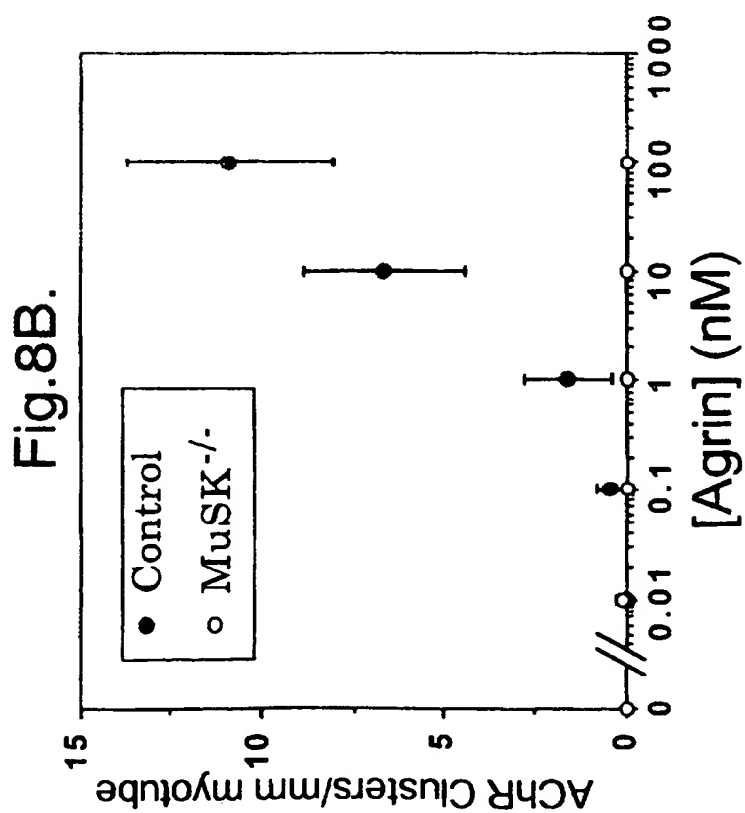


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Fig.7C.



MuSK -/-



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Fig.9A.

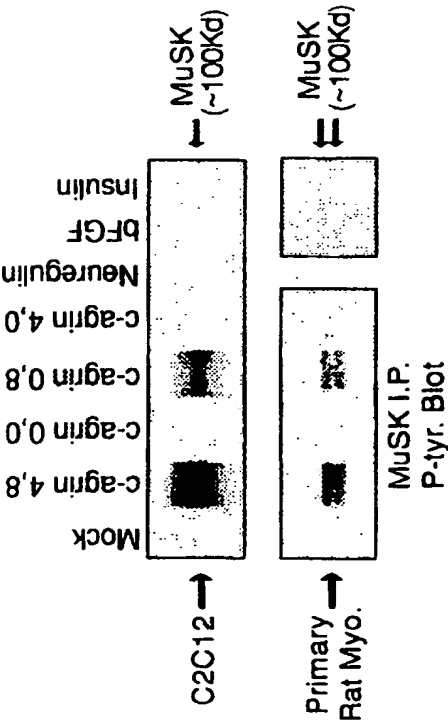


Fig.9B.

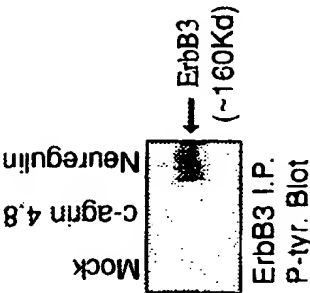


Fig.9C.

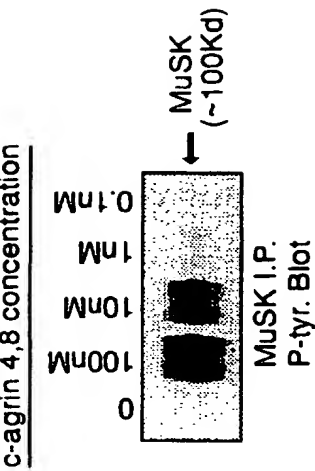


Fig.9D.



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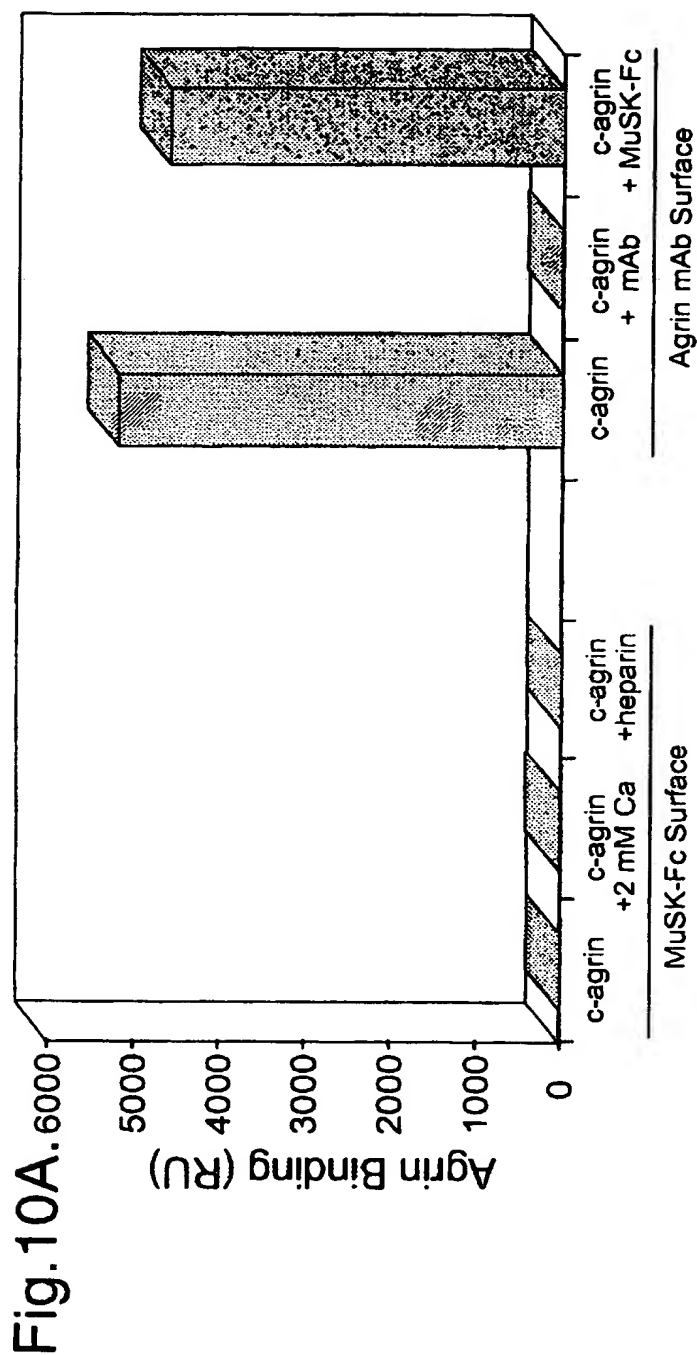
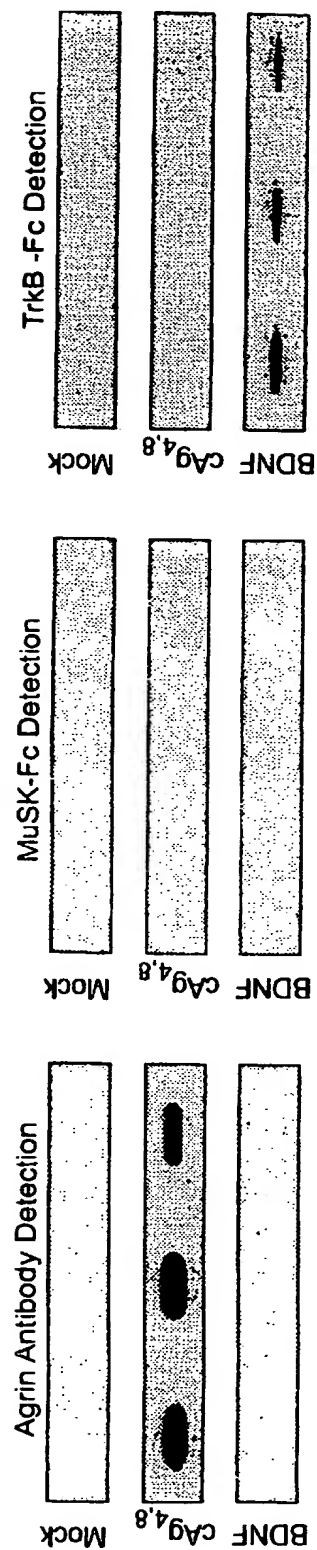


Fig. 10B.



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Fig.11.

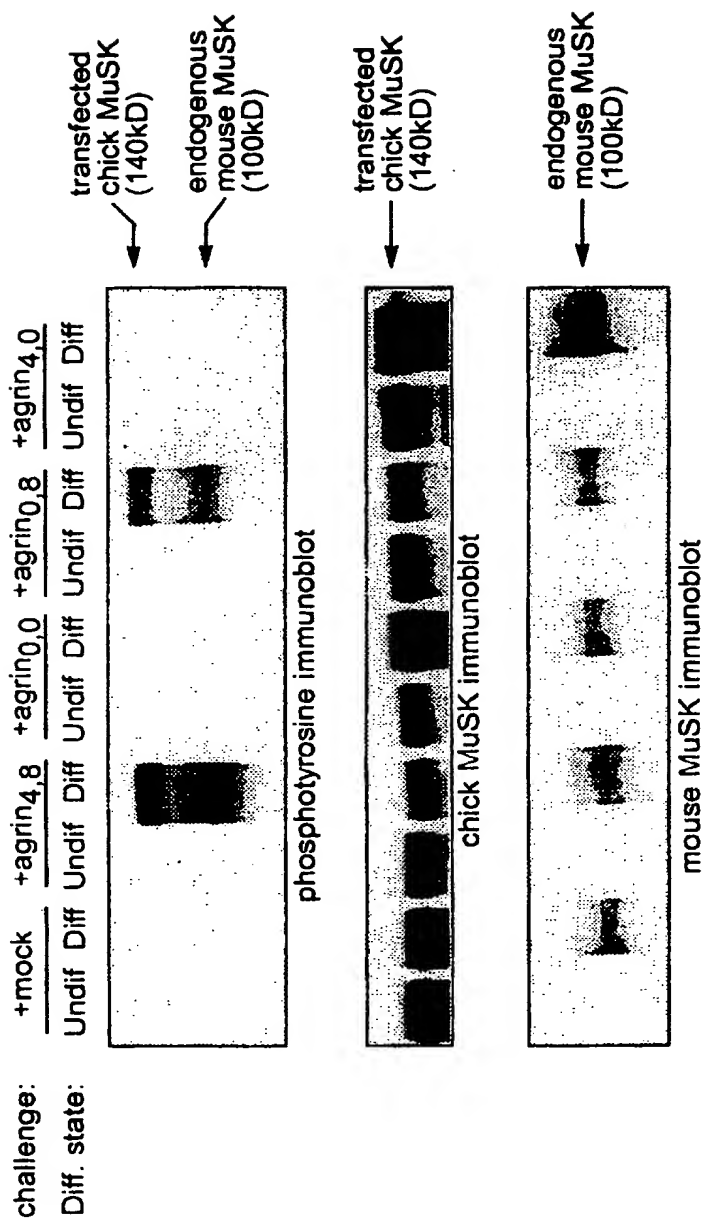
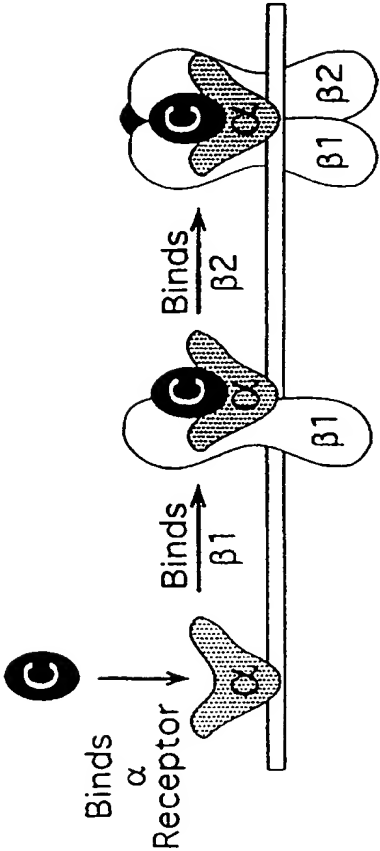
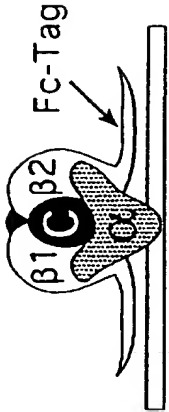


Fig.12.

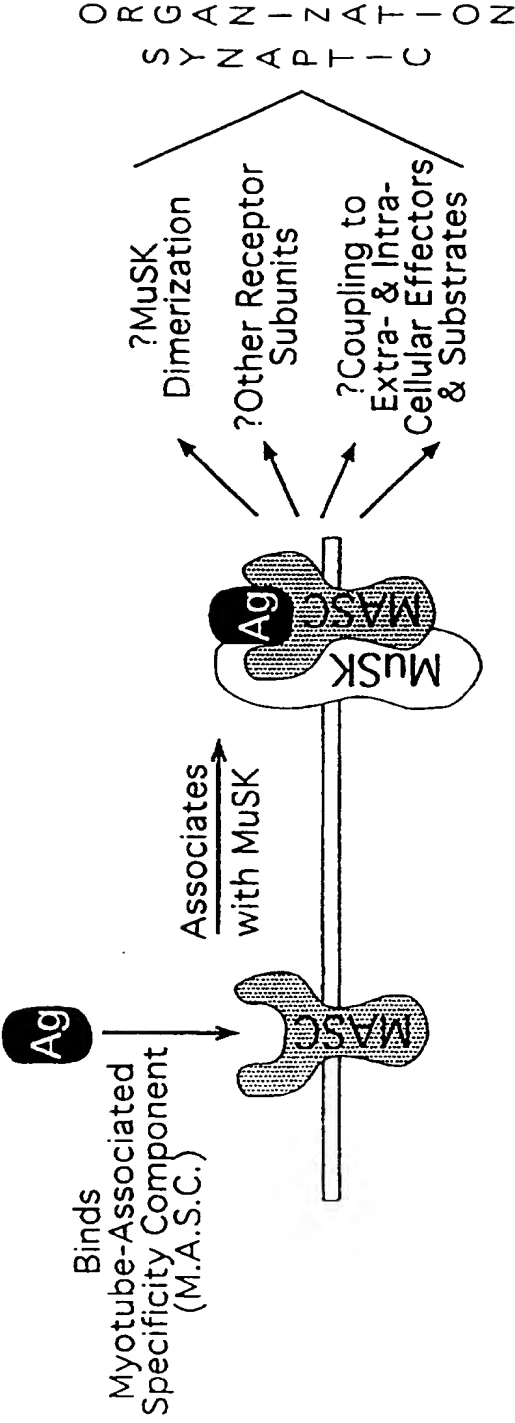
A. Formation of CNTF Receptor Complex



B. CNTF Receptor Complex With Soluble β Components



C. Formation of Receptor Complex For Agrin



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Fig. 13A.

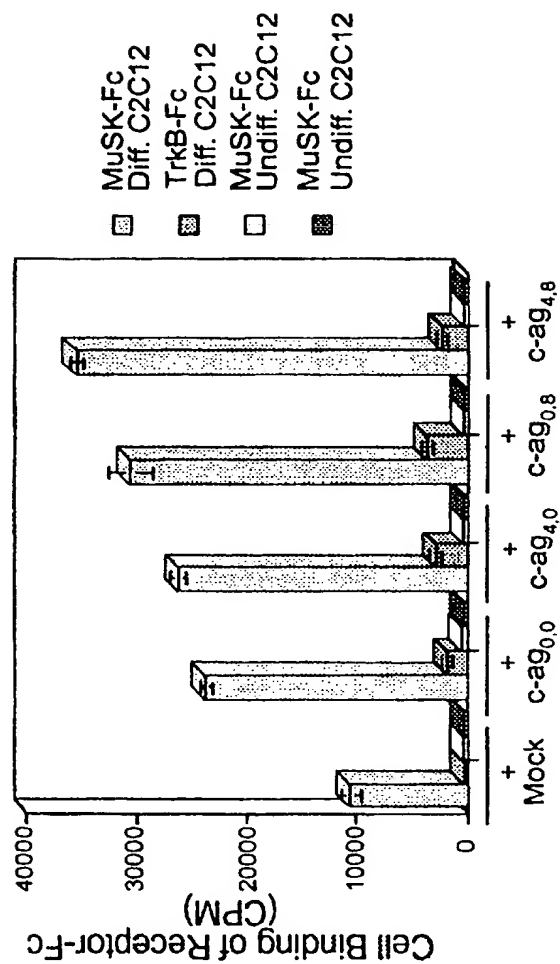


Fig. 13B.

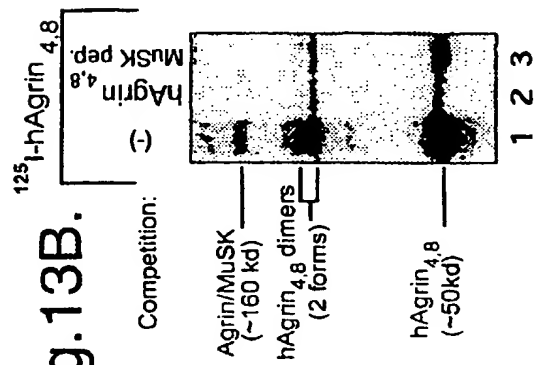
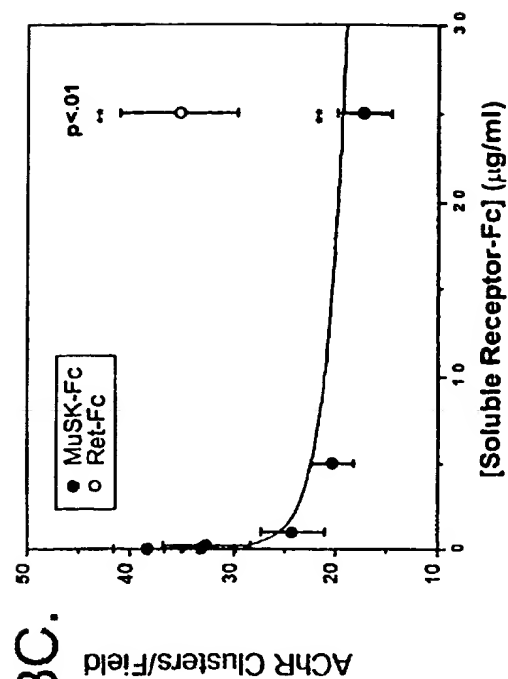


Fig. 13C.



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Fig.14.

```

      10      20      30      40      50      60
      *      *      *      *      *      *
MPPLPLEHRP RQEPGASMLV RYFMIPCNIC LILLATSTLG FAVLLFLSNY KPGIHFTPAP

      70      80      90     100     110     120
      *      *      *      *      *      *
PTPPDVCRGM LCGFGAVCEP SVEDPGRASC VCKKNACPAT VAPVCGSDAS TYSNECELQR

      130     140     150     160     170     180
      *      *      *      *      *      *
AQCNQQRIR LLRQGPGSR DPCANVTCSF GSTCVPSADG QTASCLCPTT CFGAPDGTVC

      190     200     210     220     230     240
      *      *      *      *      *      *
GSDGVDYPSE CQLLSHACAS QEHIFFKFNQ PCDPCQGSMS DLNHICRVNP RTRHPEMLLR

      250     260     270     280     290     300
      *      *      *      *      *      *
PENCPAQHTP ICGDDGVITY NDCVMSRIGA TRGLLLQKVR SGQCQTRDQC PETCQFNQVC

      310     320     330     340     350     360
      *      *      *      *      *      *
LSRRGRPHCS CDRVTCDSY RPYCAQDQHT YNNDQWRQQA ECRQQRAPP KHQGPCDQTP

      370     380     390     400     410     420
      *      *      *      *      *      *
SPCHGVQCAF GAVCTVKNQK AECECQRVCS GIYDPVCGSD GVTYGSVCEL ESMACTLQRE

      430     440     450     460     470     480
      *      *      *      *      *      *
IQVARRGPCD PCGQCRFGSL CEVETGRCVC PSECVEAQP VCGSDGHTYA SECELHVHAC

      490     500     510     520     530     540
      *      *      *      *      *      *
THQISLYVAS AGHCQTCGEK VCTFGAVCSA GQCVCPCEH PPPGPVCGSD GVTYLSACEL

      550     560     570     580     590     600
      *      *      *      *      *      *
REAAQQQVQ IEEAHAGPCE PAECGSGGSG SGEDDECEQE LCRQRGGIWD EDSQDGPVCV

      610     620     630     640     650     660
      *      *      *      *      *      *
DFSCQSVPRS PTCGSDGVITY GTECDLKKAR CESQQELYVA AQGACRGPTL APLLPVAFPH

      670     680     690     700     710     720
      *      *      *      *      *      *
CAQTPYGCCQ DNFTAAQGVG LAGCPSTCHC NPHGSYSGTC DPATGQCSCR PGVGGRLRCR

      730     740     750     760     770     780
      *      *      *      *      *      *
CEPGFVNFRG IVDGHSQCT PCSCDPRGAV RDDCEQMTGL CSCRPGVAGP KCGQCPDQVQ

      790     800     810     820     830     840
      *      *      *      *      *      *
LGHLCQADP MTPVTCVEIH CEFGASCVEK AGFAQCICPT LTCPEANSTK VCGSDGVITY

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Fig.14 (Cont 1).

850	860	870	880	890	900
* *	* *	* *	* *	* *	* *
NECQLKAIAC	RQRLDISTQS	LGPCQESVTP	GASPTSASMT	TPRHILSKTL	PFPHNSLPLS
910	920	930	940	950	960
* *	* *	* *	* *	* *	* *
PGSTTHDWPT	PLPISPHTTV	SIPRSTAWPV	LTVPPTAAAS	DVTSLATISF	SESGSANGSG
970	980	990	1000	1010	1020
* *	* *	* *	* *	* *	* *
DEELSGDEEA	SGGGSGGLEP	FVGSIVVTHG	PPIERASCYN	SPLGCCSDGK	TPSLDSEGSN
1030	1040	1050	1060	1070	1080
* *	* *	* *	* *	* *	* *
CPATKAFQGV	LELEGVEGQE	LFYTPEMADP	KSELFGETAR	SIESTLDDLF	RNSDVKKDFW
1090	1100	1110	1120	1130	1140
* *	* *	* *	* *	* *	* *
SVRLRELPG	KLVRIVDVH	FDPTTAFQAS	DVGQALLRQI	QVSRPWALAV	RRPLQEHVRF
1150	1160	1170	1180	1190	1200
* *	* *	* *	* *	* *	* *
LDFDWFPTFF	TGAATGTAA	MATARATTVS	RLPASSVTPR	VYPSHTSRPV	GRTTAPPTTR
1210	1220	1230	1240	1250	1260
* *	* *	* *	* *	* *	* *
RPPTTATNMD	RPRTPGHQQP	SKSCDSQPCL	HGGTCQDQDS	GKGFTCSCTA	GRGGSVCEKV
1270	1280	1290	1300	1310	1320
* *	* *	* *	* *	* *	* *
QPPSMPAFKG	HSFLAFPTLR	AYHTLRLALE	FRALETEGLL	LYNGNARGKD	FLALALLDGR
1330	1340	1350	1360	1370	1380
* *	* *	* *	* *	* *	* *
VQFRFDTGSG	PAVLTSVLPV	EPGRWHRLEL	SRHWRQGTLS	VDGETPVVGE	SPSGTDGLNL
1390	1400	1410	1420	1430	1440
* *	* *	* *	* *	* *	* *
DTNLYVGGIP	EEQVAMVLDL	TSVGVGLKGC	IRMLDINNQQ	LELSDWQRAA	VQSSGVGECG
1450	1460	1470	1480	1490	1500
* *	* *	* *	* *	* *	* *
DHPCLPNPCH	GGALCQALEA	GMFLCQCPPG	RFGPTCADEK	SPCQPNPCHG	AAPCRVLSSG
1510	1520	1530	1540	1550	1560
* *	* *	* *	* *	* *	* *
GAKCECPLGR	SGTFCQTVLE	TAGSRPFLLD	FNGFSYLELK	GLHTFERDLG	EKMALEMVFL
1570	1580	1590	1600	1610	1620
* *	* *	* *	* *	* *	* *
ARGPSGLLLY	NGQKTDGKGD	FVSLALHNRH	LEFCYDLGKG	AAVIRKEPI	ALGTWVRVFL
1630	1640	1650	1660	1670	1680
* *	* *	* *	* *	* *	* *
ERNGRKGALQ	VGDGPRVLGE	SPKSRKVPHT	MLNLKEPLYI	GGAPDFSKLA	RGAUVSSGFS

▲
Y-site

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Fig.14 (Cont 2).

1690	1700	1710	1720	1730	1740
* *	* *	* *	* *	* *	* *
GVIQLVSLRG	HQLLTQEHVL	RAVDVSPFAD	HPCTQALGNP	CLNGGSCVPR	EATYECLCPG
1750	1760	1770	1780	1790	1800
* *	* *	* *	* *	* *	* *
GFSGLHCEKG	LVEKSVGDL	TLAFDGRTYI	EYLNNAVIESE	KALQSNHFEL	SLRTEATQGL
			▲ Z-site		
1810	1820	1830	1840	1850	1860
* *	* *	* *	* *	* *	* *
VLWIGKAAER	ADYMALAIVD	GHLQLSYDLG	SQPVVLRSTV	KVNTNRWLRI	RAHREHREGS
1870	1880	1890	1900	1910	1920
* *	* *	* *	* *	* *	* *
LQVGNEAPVT	GSSPLGATQL	DTDGALWLGG	LQKLPVGQAL	PKAYGTGFVG	CLRDVVVGHR
1930	1940				
* *	* *				
QLHLLEDAVT	KPELRPCPTP	*			

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Fig.15.

ATG	TCT	GCA	CTT	CTG	ATC	CTA	GCT	CTT	GTT	GGA	GCT	GCA	GTT	GCT	GAC
M	S	A	L	L	I	L	A	L	V	G	A	A	V	A	D
TAC	AAA	GAC	GAT	GAC	GAC	AAG	AAG	AGC	CCC	TGC	CAG	CCC	AAC	CCC	TGC
Y	K	D	D	D	D	K	K	S	P	C	Q	P	N	P	C
CAT	GGG	GCG	GCG	CCC	TGC	CGT	GTG	CTG	CCC	GAG	GGT	GGT	GCT	CAG	TGC
H	G	A	A	P	C	R	V	L	P	E	G	G	A	Q	C
GAG	TGC	CCC	CTG	GGG	CGT	GAG	GGC	ACC	TTC	TGC	CAG	ACA	GCC	TCG	GGG
E	C	P	L	G	R	E	G	T	F	C	Q	T	A	S	G
CAG	GAC	GGC	TCT	GGG	CCC	TTC	CTG	GCT	GAC	TTC	AAC	GGC	TTC	TCC	CAC
Q	D	G	S	G	P	F	L	A	D	F	N	G	F	S	H
CTG	GAG	CTG	AGA	GGC	CTG	CAC	ACC	TTT	GCA	CGG	GAC	CTG	GGG	GAG	AAG
L	E	L	R	G	L	H	T	F	A	R	D	L	G	E	K
ATG	GCG	CTG	GAG	GTC	GTG	TTC	CTG	GCA	CGA	GGC	CCC	AGC	GGC	CTC	CTG
M	A	L	E	V	V	F	L	A	R	G	P	S	G	L	L
CTC	TAC	AAC	GGG	CAG	AAG	ACG	GAC	GGC	AAG	GGG	GAC	TTC	GTG	TCG	CTG
L	Y	N	G	Q	K	T	D	G	K	G	D	F	V	S	L
GCA	CTG	CGG	GAC	CGC	CGC	CTG	GAG	TTC	CGC	TAC	GAC	CTG	GGC	AAG	GGG
A	L	R	D	R	R	L	E	F	R	Y	D	L	G	K	G
GCA	GCG	GTC	ATC	AGG	AGC	AGG	GAG	CCA	GTC	ACC	CTG	GGA	GCC	TGG	ACC
A	A	V	I	R	S	R	E	P	V	T	L	G	A	W	T
AGG	GTC	TCA	CTG	GAG	CGA	AAC	GGC	CGC	AAG	GGT	GCC	CTG	CGT	GTG	GGC
R	V	S	L	E	R	N	G	R	K	G	A	L	R	V	G
GAC	GGC	CCC	CGT	GTG	TTG	GGG	GAG	TCC	CCG	AAA	TCC	CGC	AAG	GTT	CCG
D	G	P	R	V	L	G	E	S	P	K	S	R	K	V	P
CAC	ACC	GTC	CTC	AAC	CTG	AAG	GAG	CCG	CTC	TAC	GTA	GGG	GGC	GCT	CCC
H	T	V	L	N	L	K	E	P	L	Y	V	G	G	A	P
GAC	TTC	AGC	AAG	CTG	GCC	CGT	GCT	GCT	GCC	GTG	TCC	TCT	GGC	TTC	GAC
D	F	S	K	L	A	R	A	A	A	V	S	S	G	F	D
GGC	GCC	ATC	CAG	CTG	GTC	TCC	CTC	GGA	GGC	CGC	CAG	CTG	CTG	ACC	CCG
G	A	I	Q	L	V	S	L	G	G	R	Q	L	L	T	P
GAG	CAC	GTG	CTG	CGG	CAG	GTG	GAC	GTC	ACG	TCC	TTT	GCA	GGT	CAC	CCC
E	H	V	L	R	Q	V	D	V	T	S	F	A	G	H	P
TGC	ACC	CGG	GCC	TCA	GGC	CAC	CCC	TGC	CTC	AAT	GGG	GCC	TCC	TGC	GTC
C	T	R	A	S	G	H	P	C	L	N	G	A	S	C	V
CCG	AGG	GAG	GCT	GCC	TAT	GTG	TGC	CTG	TGT	CCC	GGG	GGA	TTC	TCA	GGA
P	R	E	A	A	Y	V	C	L	C	P	G	G	F	S	G
CCG	CAC	TGC	GAG	AAG	GGG	CTG	GTG	GAG	AAG	TCA	GCG	GGG	GAC	GTG	GAT
P	H	C	E	K	G	L	V	E	K	S	A	G	D	V	D

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Fig.15 (Cont).

ACC	TTG	GCC	TTT	GAC	GGG	CGG	ACC	TTT	GTC	GAG	TAC	CTC	AAC	GCT	GTG
T	L	A	F	D	G	R	T	F	V	E	Y	L	N	A	V
ACC	GAG	AGC	GAA	CTG	GCC	AAT	GAG	ATC	CCC	GTC	Z-insert				
T	E	S	E	L	A	N	E	I	P	V	E	K	A	L	Q
AGC	AAC	CAC	TTT	GAA	CTG	AGC	CTG	CGC	ACT	GAG	GCC	ACG	CAG	GGG	CTG
S	N	H	F	E	L	S	L	R	T	E	A	T	Q	G	L
GTG	CTC	TGG	AGT	GGC	AAG	GCC	ACG	GAG	CGG	GCA	GAC	TAT	GTG	GCA	CTG
V	L	W	S	G	K	A	T	E	R	A	D	Y	V	A	L
GCC	ATT	GTG	GAC	GGG	CAC	CTG	CAA	CTG	AGC	TAC	AAC	CTG	GGC	TCC	CAG
A	I	V	D	G	H	L	Q	L	S	Y	N	L	G	S	Q
CCC	GTG	GTG	CTG	CGT	TCC	ACC	GTG	CCC	GTC	AAC	ACC	AAC	CGC	TGG	TTG
P	V	V	L	R	S	T	V	P	V	N	T	N	R	W	L
CGG	GTC	GTG	GCA	CAT	AGG	GAG	CAG	AGG	GAA	GGT	TCC	CTG	CAG	GTG	GGC
R	V	V	A	H	R	E	Q	R	E	G	S	L	Q	V	G
AAT	GAG	GCC	CCT	GTG	ACC	GGC	TCC	TCC	CCG	CTG	GGC	GCC	ACG	CAG	CTG
N	E	A	P	V	T	G	S	S	P	L	G	A	T	Q	L
GAC	ACT	GAT	GGA	GCC	CTG	TGG	CTT	GGG	GGC	CTG	CCG	GAG	CTG	CCC	GTG
D	T	D	G	A	L	W	L	G	G	L	P	E	L	P	V
GGC	CCA	GCA	CTG	CCC	AAG	GCC	TAC	GGC	ACA	GGC	TTT	GTG	GGC	TGC	TTG
G	P	A	L	P	K	A	Y	G	T	G	F	V	G	C	L
CGG	GAC	GTG	GTG	GTG	GGC	CGG	CAC	CCG	CTG	CAC	CTG	CTG	GAG	GAC	GCC
R	D	V	V	V	G	R	H	P	L	H	L	L	E	D	A
GTC	ACC	AAG	CCA	GAG	CTG	CGG	CCC	TGC	CCC	ACC	CCA	TGA			
V	T	K	P	E	L	R	P	C	P	T	P	*			

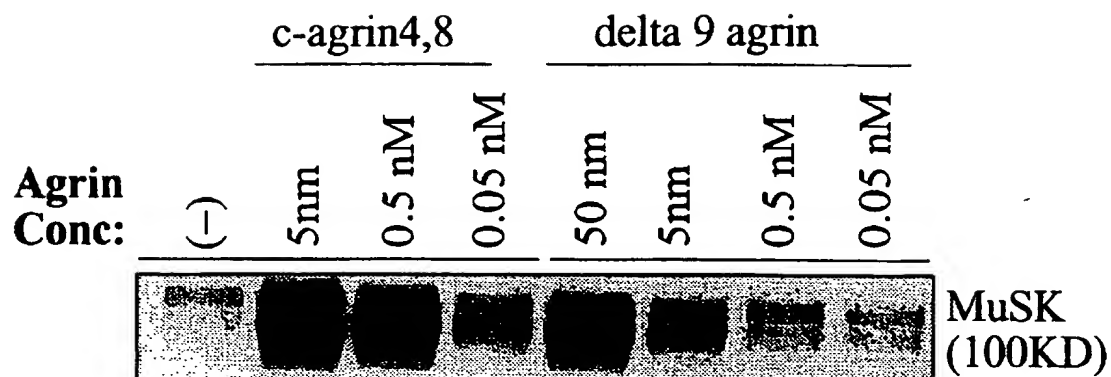
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Fig.16.

Fig.17.

